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(SA) Thiazolidine and pyrrolidine compounds, processes for their preparation and pharmaceutical compositions containing

(3) Thiazolidine and pyrrolidine compounds which have the general formula

$$R^{A} \xrightarrow{Q^{1}-Q^{2}} CO-R^{b}$$

and saits thereof for preventing or relieving diabetic complications and for reducing blood pressure, the processes for their preparation, and the compositions comprising them and pharmaceutically acceptable excipient(s).

TITLE OF INVENTION

THIAZOLIDINE AND PYRROLIDINE COMPOUNDS, PROCESSES FOR THEIR PREPARATION AND PHARMACEUTICAL COMPOSITIONS CONTAINING

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BACKGROUND OF INVENTION

This invention relates to thiazolidine and pyrrolidine compounds of the general formula

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wherein Q^1 and Q^2 each is methylene or sulfur, but Q^1 and Q^2 are not sulfur at the same time;

RA is Ra or Rb;

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RB and RC each is RC;

$$\text{W is} \quad \begin{bmatrix} R^{\frac{1}{2}} \\ C \\ R^{\frac{1}{2}} \end{bmatrix}_{\mathcal{L}} \quad \begin{bmatrix} R^{\frac{1}{2}} \\ R^{\frac{1}{2}} \end{bmatrix}_{\mathcal{L}} \quad \begin{bmatrix}$$

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x, y and Z each is single bond, $-CH_2^-$, $-C = C_-$

1 -0-,-co-, -s-, -so-, -so₂-, -c-, -NHCONH-, -N N- or -N-, |
$$\frac{1}{N-R}$$
 | $\frac{1}{N-R}$ | $\frac{1}{N-$

 R^{22} , R^{23} , R^{24} , R^{25} and R^{26} each is R^{d} ;

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R^a is selected from the group consisting of

(i) hydrogen, lower alkyl and lower alkenyl, and

(ii) lower alkyl and lower alkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, hydroxy, lower alkoxy, halogenolower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfcnyl and lower alkylsulfinyl;

R^b is selected from the group consisting of

(a) (i) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl, and

(ii) aralkyl, heteroalkyl, aralkenyl and heteroaralkenyl

substituted by at least one substituent selected from the

group consisting of lower alkyl, lower alkenyl, halogeno
lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy,

acyloxy, halogen, nitro, cyano, amino, lower alkylamino,

dialkylamino, acylamino, mercapto acylmercapto, lower

alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxy
carbonyl, aryloxycarbonyl, sulfamoyl, lower alkylamino
sulfonyl and lower alkylsulfinyl, and

(iii) carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl,

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- aryloxycarbonyl and heteroaryloxycarbonyl;
 - (b) (i) phenyl and naphthyl, and
- (ii) phenyl and naphthyl substituted by at least one substituen selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxy-carbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl,
 - (c)(i) furyl, thienyl and pyridyl, and

lower alkylaminosulfonyl and lower alkylsulfinyl;

(ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

R^C is selected from the group consisting of

(a) (i) hydroxy, lower alkoxy and amino, and

(ii) lower alkoxy, and amino substituted by at least one substituent selected from the group consisting of lower alkyl, aralkyl, heteroaralkyl, aralkenyl, heteroaralkenyl, hydroxy, lower alkoxy, aralkyloxy, heteroaralkyloxy, aryloxy, heteroaryloxy, acyloxy, aryl, heteroaryl, substituted aralkyl and substituted aryl wherein the substituent is lower alkyl, lower alkoxy, halogen or amino;

- (b)(i) aryloxy and heteroaryloxy, and
- (ii) aryloxy and heteroaryloxy substituted by at least one substituent selected from the group consisting of lower alkyl, hydroxy, lower alkoxy, halogen and amino, and

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- R^d is selected from the group consisting of

 (a) (i) hydrogen, lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy,
- carboxy, amino, mercapto and sulfo, and

 (ii) lower alkyl, lower alkenyl, aralkyl, heteroaralkyl,

 alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy,

 amino, mercapto and sulfo substituted by at least one substituent

 selected from the group consisting of lower alkyl, lower alkenyl,

 lower alkoxy, lower alkanoyl, aryl, heteroaryl, acyloxy, aroyl,

 hydroxy, carboxy, amino, guanidino, mercapto, acylamino,

 acylmercapto, lower alkoxycarbonyl, sulfo, halogen, nitro,

 cyano, sulfamoyl, lower alkylaminosulfonyl, lower alkylthio

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and lower alkylsulfinyl;

- (b) (i) phenyl and naphthyl, and

 (ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogenolower alkyl, halogenolower alkoxy, lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
- (c) (i) furyl, thienyl and pyridyl, and
 (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkvl, halogeno-lower alkoxy, lower alkylene-dioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkyl-aminosulfonyl and lower alkylsulfinyl;
- and salts thereof which are useful as agents for therapy or prophylaxis of the diabetic complication because they inhibit strongly aldose reductase, and as antihypertensive agents because they inhibit angiotensin I-converting enzyme.

The compounds [I] of this invention can be prepared by following process.

(i) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II]

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$$R^{A} \xrightarrow{Q^{1}-Q^{2}} CO-R^{B}$$
 [II],

wherein R^A and R^B may be protected by any suitable groups

(e.g., lower alky1, acyl, aralky1, aralkyloxy, etc.) when
R^A and R^B include reactive groups (e.g., amino, hydroxy,
mercapto, hydroxyamino, etc.), with the reactive derivative
of carboxylic acid of the formula [III] (e.g., acyl halide,
acid anhydide, mixed anhydride, active ester, lactone, etc.)

by general methods used in peptide syntheses or amide
formation reactions

wherein W and R^C may be protected by any suitable groups

(e.g., lower alkyl, acyl, aralkyl, aralkyloxy, etc.) when
W and R^C include reactive groups (e.g., amino, hydroxy,
mercapto, hydroxyaminc, etc.), followed by removal of
protective groups by well-known methods (e.g., hydrolysis,
hydrogenolysis, ammonclysis, alcoholysis, etc.).

This procedures of deprotection can be applied in the following methods.

(ii) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II] with the reactive derivative of carboxylic acid of [IV] (e.g., above-mentioned)

5 $HOOC-W^1-L$ [IV],

wherein W^1 is $\begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}$, and may be protected such as (i)

above, I is a leaving group (e.g., halogen, alkylsulfonyl, arylsulfonyl, etc.), by the same methods as described in (i) above to produce a compound of the formula [V]

$$R^{A} \xrightarrow{Q^{1}-Q^{2}} CO-R^{B}$$

$$[V]$$

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and then reation of a compound of the formula [V] with a compound of the formula [VI]

20 (H)
$$X-W^2-Y-W^3-Z-W^4-CO-R^C$$
 [VI],

wherein
$$W^2$$
 is
$$\begin{bmatrix} R^5 \\ C \\ C \\ R^6 \end{bmatrix}_n \begin{bmatrix} R^7 \\ R^8 \end{bmatrix}_P \quad W^3 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_r \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \quad W^4 \end{bmatrix}_{R} \quad W^4 \text{ is } \begin{bmatrix} R^9 \\ R^9 \end{bmatrix}_{R} \quad W^4 \end{bmatrix}_{R} \quad W^4$$

protected such as (i) above, in the presence of proper alkaline and/or organic bases, if necessary, by known methods.

(iii) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [VII] (e.g., metioned in (i) above)

$$HOOC-W^1-X-W^2-L$$
 [VII]

and then with a compound of the formula (VIII)

$$Y-W^3-z-W^4-CO-R^C$$
 [VIII]

by the same method as (ii) above.

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(iv) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [IX] (e.g., mentioned in (i) above)

HOOC-
$$w^1$$
-x- w^2 -y- w^3 -L [IX],

and then with a compound of the formula [X]

$$(H) Z-W^4-CO-R^C$$

by the same method as (ii) above.

[XI],

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(v) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II] with the reactive derivative of carboxylic acid [XI] (e.g., acyl halide, acid anhydride, mixed anhydride, active ester, lactone, thiolactone, etc.)

 $HOOC-W^1-X(H)$

and then with a compound of the formula [XII]

10 $L-W^2-Y-W^3-Z-W^4-CO-R^C$ [XII]

by the same method as (ii) above.

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(vi) A compound of the formula [I] is yielded by the
reaction of a compound of the formula [II] with the reactive
derivative of carboxylic acid of the formula [XIII] (e.g.,
mentioned in (v) above)

$$HOOC-W^1-X-W^2-Y(H)$$
 [XIII],

and then with a compound of the formula [XIV]

$$L-W^3-Z-W^4-CO-R^C \qquad \qquad [XIV]$$

by the same method as (ii) above.

(vii) A compound of the formula [I] is yielded by the reaction of a compound of the formula [II] with the

reactive derivative of carboxylic acid of the formula '[XV] (e.g., mentioned in (v) above)

$$HOOC-W^{1}-X-W^{2}-Y-W^{3}-Z(H)$$
 [XV],

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and then with a compound of the formula [XVI]

by the same method as (ii) above.

(viii) A compound of the formula [I] is also yielded by converting a compound of the formula [I] prepared by any method above-mentioned by well-known methods (e.g., oxidation, formation of oxime, hydrazone and semicarbazone, addition to double bond, etc.)

The compounds [I] of this invention are effective on preventing or relieving diabetic complications.

In diabetic patients, high levels of hexoses (e.g.,

glucose, galactose. etc.) in blood lead to the accumulation
of sugar alcohols (e.g., sorbitol, galactitol, etc.) in
tissues. It is known that this accumulation causes the
swelling of cells to induce complications of diabetic
cataract, diabetic retinopathy, diabetic nephropathy, diabetic
neuropathy, etc. [R. Quan-Ma et al., Biochem. Biophys. Res.
Comm., 22, 492 (1966)]. For example, R. Gitzelman et al.

have presented that cataract is caused by the accumulation of sugar alcohols [Exptl. Eye. Res., 6, 1 (1967)]. A report of Kinoshita et al. has demonstrated that aldose reductase which reduced aldose to the corresponding sugar alcohols

was involved in the initiation of these diabetic complications and that effective inhibitors of almose reductase were useful [Jpn. J. Ophthalmol., 20, 339 (1976)]. On the basis of the above information, aldose reductase inhibition of the compounds [I] of this invention was tested.

The results of the examinations demonstrated that these compounds have potent inhibitory activities on aldose reductase, and therefore they are useful as drugs for therapy or prophylaxis of the diabetic complications.

On the other hand, it has been known that a kind of the derivatives of thiazolidine- or pyriolidinecarboxylic acid have potent inhibitory activity to angiotensin I-converting enzyme, but thiazolidine and pyrrolidine compounds of this invention are novel compounds, and have more potent inhibitory activities to angiotensin I-converting enzyme. Furthermore, the compounds of this invention are prepared by convenient methods, and are superior to the stability.

Thus, the compounds of this invention are useful as therapeutic or prophylactic agents and antihypertensive agents.

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The compound of formula [I] can form the conventional

salts to be generally used as medicine such as sodium salt, potassium salt, calcium salt, magnesium salt, alminum salt, ammonium salt, diethylamine salt, triethanolamine, etc.

The compounds of formula [I] have the stereoisomers which are within the limit of this invention, because they have one or more asymmetric carbon atoms.

Typical examples are shown below, although this invention is not limited to these examples.

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EXAMPLE 1

(4R)-3-(7-Carboxyheptanoy1)-2-(2-hydroxypheny1)-4thiazolidinecarboxylic acid (compound 20)

(4R)-2-(2-Hydroxyphenyl)-4-thiazolidinecarboxylic acid 5 (6.8g,) in N sodium hydroxide (30ml) and octanedicyl dichloride (6.3g,) were added dropwise to 1M potassium carbonate (60ml) with stirring under ice-cooling. After the addition, the reaction mixture was stirred for 1 hour at the same temperature and for additional 1 hour at room 10 The solution was acidified with dilute hydrochloric acid, and extracted with ethyl acetate. organic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate, and evaporated in vacuo. The residual oil *2 was purified by 15 silica gel column chrcmatography to give 7.0g (61%) of the titled compound: mp 155-157°C (dec.) (ethyl acetate); $[\alpha]_D^{27}$ +134.1° (c=0.5, methanol). IR (nujol, cm⁻¹): 3220 (OH), 1710 (COOH), 1620 (CON), 1600 (aromatic), 1415, 1235, 1172, 950, 760. NMR (DMSO-d₆, δ): 0.53-1.73 (8H, m, -CH₂+CH₂)₄-CH₂-), 20 1.77-2.57 (4H, m. $-CH_2(CH_2)_4(CH_2^-)$, 3.63 (1H, $AB_q(A part)$, d, J=11.5, 8.5Hz, C₅^{*1}-H_A), 3.37 (1H, AB_q(B part), d, J=11.5, 6.5Hz, $C_5^{-H}_B$), 4.60 (lH, dd, J=8.5, 6.5Hz, C_4^{-H}), 6.28 (1H, s, C_2 -H), 6.45-8.07 (4H, m, arom. H), 9.77 (1H, s, -COOH). TLC: Rf value*3 0.52. 25

tl The numbers represent the positions on thiazolidine or



pyrrolidinc ring. The same shall be applied hereinafto

- *2 Two spots were observed on the TLC (ethyl acetatechloroform-acetic acid (10:5:3)), and two products could be separated by silica gel column chromatography
- From NMR spectra, the upper and lower spots were identified as the titled compound and (4R,4R')-3,3'-(octanedicyl)bis[2-(2-hydroxyphenyl)-4-thiazolidine-carboxylic acid) (compound 40), respectively.
- *3 Silica gel, ethyl acetate-chloroform-acetic acid (10:5:3).

The compounds shown in Table I and III were prepared by the same procedure as described above.

The following compounds are also prepared by the same procedure as EXAMPLE 1.

- (4R)-3-carboxyacetyl-4-thiazolidinecarboxylic acid
 (4R)-3-(3-carboxypropanoyl)-2-phenyl-4-thiazolidinecarboxylic acid
- (4R)-3-[3-(2-carboxyethylsulfinyl)propanoyl]-2-(2-
- hydroxyphenyl)-4-thiazolidinecarboxylic acid

 (4R)-3-[[[2-(carboxymethyloxy)ethyl]oxy]acetyl]-2-(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid

 (4R)-3-(4-carboxytutanoyl)-2-(3-hydroxyphenyl)-4-thiazolidinecarboxylic acid
- 25 (4R)-3-(5-carboxypentanoy1)-2-(4-methylpheny1)-4thiazolidinecarboxylic acid



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(4R)-3-(6-carboxyhexanoy1)-2-(4-chloropheny1)-4-
 <u>_</u>1
            thiazolidinecarboxylic acid
            (4R)-3-(7-carboxyheptanoy1)-2-(4-methoxypheny1)-4-
            thiazolidinecarboxylic acid
            (4R)-3-(13-carboxytridecanoy1)-2-(2-nitropheny1)-4-
  5
            thiazolidinecarboxylic acid
            (4R)-3-(7-carboxyheptanoyl)-2-(3-nitrophenyl)-4-
            thiazolidinecarboxylic acid
            (4R)-3-[3-(2-carboxyethylthio)propanoyl]-2-(5-nitro-
            phenyl) -4-thiazolidinecarboxylic acid
 10
            (4R) -3-[[[2-(carboxymethyloxy)ethyl]oxy]acetyl]-2-
            (3-nitrophenyl)-4-thiazolidinecarboxylic acid
            (4R)-3-(6-carboxyhexanoyl)-2-(4-nitrophenyl)-4-
            thiazolidinecarboxylic acid
            (4R)-3-(9-carboxynonanoy1)-2-(4-nitropheny1)-4-
15
            thiazolidinecarboxylic acid
            (4R)-3-(ll-carboxyundecanoyl)-2-(4-nitrophenyl)-4-
            thiszolidinecarboxylic acid
            (4R) -3-[4-(3-carboxypropyloxy) butanoyl]-2-(4-nitrophenyl) -
            4-thiazolidinecarboxylic acid
20
            (4R)-3-[3-(2-carboxyethylsulfonyl)propanoyl]-2-(4-nitro-
           phenyl)-4-thiazolidinecarboxylic acid
            (4R)-3-(9-carboxynonanoy1)-2-(5-chloro-2-hydroxypheny1)-
            4-thiazolidinecarboxylic acid
25
            (4R)-3-(11-carboxyundecancyl)-2-(3,4,5-trimethoxyphenyl)-
           4-thiazolidinecarboxylic acid
           (4R) -3-(13-carboxycridecanoyl) -2-(2-acetoxyphenyl) -4-
```

thiazolidinecarboxylic acid 1 (4R) -3-(6-carboxyhexanoyl) -2-(2-furyl) -4-thiazolidinecarboxylic acid (4R)-3-(7-carboxyheptanoyl)-2-(2-thienyl)-4-thiazolidine-5 carboxylic acid (4R)-3-(8-carboxyoctanoyl)-2-(3-pyridyl)-4-thiazolidinecarboxylic acid (4R) -3-(9-carboxynonanoyl) -2-(1-naphthyl) -4-thiazolidinecarboxylic acid 10 (4R)-3-(5-carboxypentanoyl)-2-(2-hydroxy-4-sulfamoylphenyl) -4-triazolidinecarboxylic acid (4R) -3-(6-carboxyhexanoyl) -2-(3-cyanophenyl) -4thiazolidinecarboxylic acid (4R) -3-(7-carboxyheptanoyl) -2-(3-difluoromethoxyphenyl) -15 4-thiazolidinecarboxylic acid (4R)-3-(8-carboxyoctanoy1)-2-(4-carboxypheny1)-4thiazolidinecarboxylic acid (4R) -3-(9-carboxynonanoy1) -2-(3-methylsulfinyiphenyl) -4thiazolidinecarboxylic acid 20

EXAMPLE 2

(4R,4'R)-3,3'-(Octanedioy1)bis[2-(2-hydroxypheny1)-4-thiazolidinecarboxylic acid (compound 40)

To a stirred solution of (4R)-2-(2-hydroxyphenyl)4-thiazolidinecarboxylic acid (6.8g) in lM
potassium carbonate (45ml), octanedicyl dichloride (3.2g)

was added dropwise under ice-cooling. After the addition, the reaction mixture was stirred for 1 hour at the same temperature and for additional 1 hour at room temperature. The solution was acidified with dilute hydrochloric acid, extracted with ethyl acetate. The

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organic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate, and evaporated in vacuo. The residual oil was purified by silica gel column chromatography to give 7.6g (86%) of the titled compound: mp 93-97°C (dec.); [a] 1/2 +123.6° (c=0.5, methanol). IR (nujol, cm⁻¹): 1720 (COOH), 1620 (CON), 1600 (aromatic), 1230, 1090, 855, 765. MNR (CD3OD) 6: 0.7-1.7 (8H, m, -CH2 (CH2) - CH2), 1.8-2.4 (4H, m, -CH2 (CH2) - CH2), 3.25 (4H, d, J=7.5Hz, C5-H), 4.81 (2H, t, J=7.5Hz, C4-H), 6.35 (2H, s, C2-H), 6.7-8.0 (8H, m, arom. H). TLC: Rf value 0.34.

* Silica gel, ethyl acetate-chloroform-acetic acid (10:5:3).

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The compounds shown in Table II and III were prepared by the same procedure as described above.

EXAMPLE 3

20 (4R,4'R)-3,3'-(heptanedioyl)bis[2-(3-cyanophenyl)-4thiazolidinecarboxylic acid] (compound 36)

To a stirred solution of (4R)-2-(3-cyanophenyl)-4thiazolidinecarboxylic acid (4.7g) in lM sodium

carbonate (30ml), heptanedicyl dichloride (2.1g)
was added dropwise under ice-cooling. The reaction mixtur
was stirred for 30 minutes at the same temperature, and

- filtered to give the precipitates. The precipitates were
 dissolved in hot water (100ml), and acidified with
 concentrated hydrochloric acid. The separated crystals
 were collected by filtration to give 3.5g (59%) of the

 titled compound: mp 105-112°C; [α]_D²⁵ +115.0° (c=1.0,
 methanol). IR (nujol, cm⁻¹): 2270 (CN), 1735 (COOH),
 1640 (CON), 1610 (arematic), 1195, 790 (arematic). NMR
 (DMSO-d₆) δ: 0.69-1.66 (6H, m, -CH₂+CH₂)₃ CH₂-),
 1.70-2.50(4H, m, -CH₂+CH₂)₃ CH₂-), 2.85-3.66 (4H, m,

 C₅-H), 4.69 (LH, dd, J=8.2, 6.0Hz, C₄-H), 5.13(1H, m;
 C₄-H), 6.16 (1H, s, C₂-H), 6.43 (1H, s, C₂-H), 7.3-8.3
 (8H, m, arom. H). TLC: Rf value 0.33.
- * Silica gel, ethyl acetate-chloroform-acetic acid (10:5:3).

The compounds shown in Table II were prepared by the same procedure as described above.

The following compounds are also prepared by the same 20 procedure as EXAMPLE 2 or 3.

- (4R,4'R)-3,3'-ipropanedicyl)bis(4-thiazolidinecarboxylic acid)
- (4R,4'R)-3,3'-(butanedioyl)bis(2-phenyl)-4-thiazolidine-carboxylic.acid)
- 25 (4R,4'R)-3,3'-(3,3!-sulfinyldipropanoyl)bis[2-(2-hydroxy-phenyl)-4-thiazolidinecarboxylic acid]
 - (4R,4'R)-3,3'-[(ethylenedioxy)diacetyl]bis[2-(2-hydroxy-phenyl)-4-thiazolidinecarboxylic acid]
 - (4R,4'R)-3,3'-[(ethylenedithio)diacetyl]bis{2-(2-hydroxy-phenyl)-4-thiazolidinecarboxylic acid}

- 5 (4R,4'R)-3,3'-(heptanedioyl)bis[2-(4-chlorophenyl)-4-thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(octanedioyl)bis[2-(4-methoxyphenyl)-4-thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(cetradecanedioyl)bis[2-(2-nitrophenyl)-4-

10 thiazolidinecarboxylic acid]

thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(3,3'-thiodipropanoyl)bis[2-(3-nitrophenyl)-

4-thiazolidinesarboxylic acid]

(4R,4!R)-3,3'-[(ethylenedioxy)diacetyl]bis[2-(3-nitrophenyl)

4-thiazolidinecarboxylic acid]

15 (4R,4'R)-3,3'-(heptanedioyl)bis[2-(4-nitrophenyl)-4-thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(decanedicyl)bis[2-(4-nitrophenyl)-4-thiazolidinecarboxylic acid]

(4R, 4'R)-3,3'-(dodecanedioyl)bis[2-(4-nitrophenyl)-4-

20 thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(4,4'-oxydibutanoyl)bis[2-(4-nitrophenyl)-

4-thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(3,3'-sulfonyldipropanoyl)bis[2-(4-nitro-phenyl)-4-thiazolidinecarboxylic acid]

25 (4R,4'R)-3,3'-(decanedicyl)bis{2-(5-chloro-2-hydroxyphenyl)-4-thiazolidinecarboxylic acid]

(4R,4'R)-3,3'-(dodecanedicyl)bis{2-(3,4,5-trimethoxy-phenyl)-4-thiazolidinecarboxylic acid}

(4R, 4R) - 3, 3 - (nonanedioyl) bis [2 - (4 - carboxyphenyl) - 4 - (4 - carboxyphenyl)]

(4R,4'R)-3,3'-(decanedioy1)bis[2-(3-methylsulfinyl-

EXAMPLE 4

(4R,4'R)-3,3'-(Heptanedioyl)bis[2-(3-nitrophenyl)-4
thiazolidinecarboxylic acid] (compound 35)

phenyl) -4-thiazolidinecarboxylic acid]

thiazolidinecarboxylic acid]

15

7

To a stirred solution of (4R)-2-(3-nitrophenyl)
25 4-thiazolidinecarboxylic acid (5.lg)inllM

sodium carbonate (40ml), heptanedioyl dichloride (2.lg)

was added dropwise under ice-cooling. The

reaction mixture was stirred for 1 hour at the same temperature, and the separated crystals were filtered to give 4.7g (69%) of the titled compound as disodium salt: mp 111-113°C (dec.) (water); [a]²⁵_D +88.2° (c=0.5, methanol) IR (nujol, cm⁻¹): 1635 (CON), 1585 (COO⁻), 1520 and 1355 (NO₂), 1095, 730. TLC: Rf value 0.28.

* Silica gel, ethyl acetate-chloroform-acetic acid (10:5:3).

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EXAMPLE 5

(4R)-3-(3-Carboxypropanoyl)-2-(2-hydroxyphenyl)-4thiazolidinecarboxylic acid (compound 6)

15 To a stirred solution of (4R)-2-(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid (4.5g) and triethylamine (4.0g) in acetone (100ml), succinic anhydride (2.0g) was added at room temperature, and stirred for 3 hours at the same temperature. The reaction mixture was concentrated 20 in vacuo, and acidified with dilute hydrochloric acid. The separated oil was extracted with ethyl acetate, and the crganic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate, and 25 evaporated in vacuo to give 4.9g (75%) of the titled compound: mp 190-191°C (dec.) (ethyl acetate-methanol); $[\alpha]_{D}^{27}$ +181.6° (c=1.0, methanol). IR (nujol, cm⁻¹): 3210

- 1 (OH), 1720 (COOH), 1618 (CON), 1602 (aromatic), 1245, 1173, 940, 763. NMR (DMSC-d₆,δ): 2.0-2.7 (4H, m, -CH₂CH₂-), 3.03 (1H, AB_q(A part), d, J=11.0, 10.0Hz, C₅-H_A), 3.36 (1H, AB_q(B part), d, J=11.0, 7.0Hz, C₅-H_B), 4.61 and 5.07 (1H, dd, J=10.0, 7.0Hz and m, C₄-H), 6.36 (1H, s, C₂-H), 6.5-8.0 (4H, arom. H). TLC: Rf value 0.35.
 - Silica gel, ethyl acetate-chloroform-acetic acid (10:5:3).
- The compounds shown in Table I and III were prepared by the same procedure as described above. The following compounds are also prepared by the same procedure as EXAMPLE 5.

 (4R)-3-(4-carboxy-4-oxobutanoy1)-2-(2-hydroxypheny1)-4-thiazolidinecarboxylic acid
- 20 EXAMPLE 6

 (4R)-3-[3-(Methoxycarbonyl)-2-methylpropanoyl]-2-(2hydroxyphenyl)-4-thiazolidinecarboxylic acid (compound 4)

To a stirred solution of (4R)-2-(2-hydroxyphenyl)
25 4-thiazolidinecarboxylic acid (11.3g) in lM sodium

carbonate (80mJ), dl-3-methoxycarbonyl-2-methylpropancyl

chloride (3.2g) was added dropwise under ice-cooling.

After the addition, the reaction mixture was stirred for 1.5 hours at the same temperature. After the filtration of solution, the filtrate was acidified with dilute hydrochloric acid, and extracted with ethyl acetate. The organic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate, and evaporated in vacuo. The residual oil was purified by silica gel column chromatography to give 7.8g (44%) of the titled compound: [\alpha]_D^{25} +161.6° (c=1.0, methanol).

IR (KBr, cm⁻¹): 3380 (OH), 1723 (COOH, COOCH₃), 1624 (CON), 1235, 1200, 1174, 764.

The compounds shown in Table I and II were prepared by the same procedure as described above.

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EXAMPLE 7

(4R) -3-(3-Carboxy-2-methylpropanoy1) -2-(2-hydroxypheny1) -4-thiazolidinecarboxylic acid (compound 5)

(4R)-3-[3-(Methoxycarbonyl)-2-methylpropanoyl]-2(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid (compound
4) (7.1g) was dissolved in 2N sodium hydroxide (40ml)
and stirred for 1 hour at room temperature. The
resulting solution was acidified with dilute hydrochloric
acid and the separated crystals were filtered to give
5.1g (75%) of the titled compound: mp 163-164°C (dec.)

- 1 (ethyl acetate); $[\alpha]_D^{25}$ +174.1° (c=1.0, methanol). IR (nujol, cm⁻¹): 3330 (OH), 1730 and 1710 (COOH), 1629 (CON), 1280, 1234, 856, 771.
 - The compounds shown in Table I and II were prepared by the same procedure as described above. The following compounds are also prepared by the same procedure as EXAMPLE 6 and 7.
- (4R)-3-[4-(carboxymethyl)benzoyl]-2-(2-hydroxyphenyl)
 4-thiazolidinecarboxylic acid
 - (4R) -3-[(4-carboxyphenyl)acetyl]-2-phenyl-4-thiazolidine-carboxylic acid
 - (4R)-3-(4-carboxy-3-butenoy1)-2-(2-hydroxypheny1)-4-thiazolidinecarboxylic acid
- (4R)-3-(4-carboxy-2-butenoy1)-2-(2-hydroxypheny1)-4thiazolidinecarboxylic acid
 - (4R)-3-(4-carboxy-3-butynoy1)-2-(2-hydroxypheny1)-4-thiazolidinecarboxylic acid
- 20 EXAMPLE 8
 - (4R)-3-[3-(N-Hydroxycarbamoyl)propanoyl]-2-(2-hydroxy-phenyl)-4-thiazolidinecarboxylic acid ethyl ester (compound 10a)
- To a stirred solution of (4R)-3-(3-carboxypropanoy1)2-(2-hydroxypheny1)-4-thiazolidinecarboxylic acid ethyl
 ester (compound 9a) (1.06g) and N-methylmorpholine (0.33ml)

- in 20ml of anhydrous tetrahydrofuran, isobutyl chloroformate (0.39ml) was added dropwise at -15°C, and stirred for additional 2 hours at this temperature. To this solution, the methanol solution of hydroxylamine (0.3g)
- was added dropwise at -50°C. The reaction mixture was stirred for 1 hour at room temperature, acidified with N hydrochloric acid, and extracted

with ethyl acetate. The organic layer was washed with saturated sodium chloride solution, dried over anhydrous

magnesium sulfate, and concentrated in vacuo. The residual oil was purified by silica gel column chlromatography to give 0.7g (63%) of the titled compound. IR (KBr, cm⁻¹) 3220, 1727, 1625, 1595, 1200, 1092, 753.

NMR (acetone-d₆, δ): 1.24 (3H, t, J=7.5Hz,, CO₂CH₂CH₃), 2.17-3.07 (4H, m, CO-(CH₂)₂CO), 3.30 (1H, AB_q(A part), d, J=10.0, 2.0Hz, C₅-H_A), 3.47 (1H, AB_q(B part), d, J=10.0, 7.0Hz, C₅-H_B), 4.14 (2H, q, J=7.5Hz, CO₂CH₂), 5.18 (1H, dd, J=2.0, 7.0Hz, C₄-H), 6.40 (1H, S, C₂-H), 6.88-7.27 (4H, m, arom. H), 8.60 (2H, br. s, NHOH), 9.77 (1H, br. s,

20 OH)

25

The compounds shown in Table I were prepared by the same procedure as described above.

EXAMPLE 9

(4R,4'R)-3,3'-(Nonanedioyl bis(2-(3-nitrophenyl)-4thiazolidinecarboxylic acid methyl ester) (compound 46) To a stirred solution of (4R,4'R)-3,3'-(nonanedioyl)bis[2-(3-nitrophenyl)-4-thiazolidinecarboxylic acid]
(compound 47) (3.3g) in ethyl acetate (50ml), 2% ether
solution of diazomethane was added dropwise until the

yellow color of diazomethane was not disappeared, and
stirred continuously for 30 minutes. The reaction mixture
was concentrated in vacuo to give 3.3g (96%) of the titled
compound: mp 61-63°C (benzene); [a]²³_D +79.4° (c=1.0,
methanol). IR (KBr, cm⁻¹): 1740, 1660, 1530, 1350,

EXAMPLE 10

(4R)-3-[(2-Carboxymethylthio-3-phenyl)propanoyl]-4-thiazolidinecarboxylic acid (compound 75a and 75b)

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(4R)-3-[(2-Merczpto-3-phenyl)propanoyl]-4-thizolidine-carboxylic acid (1.0g), potassium carbonate (0.7g), chloroacetic acid (0.2g) and potassium iodide (0.05g) were dissolved in water (5ml), and stirred for 6 hours at room temperature. The reaction mixture was acidified with 5N hydrochloric acid and extracted with ethyl acetate. The organic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate and concentrated in vacuo. The titled compounds (75a and 75b) were separated from the oily residue by silica gel column chromatography.

1	75a	75b
	yield 0.4g (37%)	0.5g (47%)
	$[\alpha]_{D}^{25}$ -52.2° (c=1.2, MeOH)	-60.4° (c=1.0, MeOH)
5	IR 1720, 1620, 1422, (neat, 1217, 756	1722, 1620, 1420, 1215, 755
10	NMR 2.67-3.63 (6H, m, (CDCl ₃ ,δ)-S-CH ₂ -CO ₂ H, C ₅ -H, -CH ₂ -Ph), 3.83-4.83 (3H, m, -CO-CH-S-, C ₂ -H), 4.98 (1H, dd, J=4.5, 6.5Hz, C ₄ -H), 7.22 (5H, s, -C ₆ H ₅) 9.55 (-CO ₂ H)	2.70-3.50 (6H, m, -S-CH ₂ -CO ₂ H, C ₅ -H, -CH ₂ -Ph), 4.00-4.57 (3H, m, -CO-CH-S-, C ₂ -H) 5.02 (1H, dd, J=4.5, 9.5Hz, C ₄ -H), - 7.23 (5H, s, -C ₆ H ₅), 10.00 (-CO ₂ H)

The compounds shown in Table IV were prepared by the same procedure as described above.

EXAMPLE 11

(4R)-3-[(Carboxymethylamino)acetyl]-2-(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid (compound 81)

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(4R)-3-Chloroacetyl-2-(2-hydroxyphenyl)-4-thiazolidine-carboxylic acid (6g) was added to a stirred solution of glycine (1.5g) in N sodium hydroxide (80ml), and stirred overnight at room temperature. The solution was adjusted to pH 1.5 by 20% hydrochloric acid and washed with ethyl acetate. The aqueous layer was adjusted to pH 3.2, and

- the separated crystals were collected by filtration to 3.28g (48.2%) of the titled compound: mp 181-182°C (dec.) (water); [α]_D²⁴ +271.2° (c=0.5, MeOH). IR (KBr, cm⁻¹): 3400, 3200, 1740, 1672, 1560, 1440, 1380, 1335, 1212, 752, 648, NMR (K₂CO₃ in D₂O, δ): 3.0-4.3 (6H, m, C₅-H, COCH₂NHCH₂CO₂H), 6.33 and 6.43 (1H, each s, C₂-H), 6.6-7.3 (3H, m, arom. H), 7.82 (1H, br. d, J=8Hz, arom. H), 9.0-10.3 (2H, br. s, -OH, -CO₂H).
- The compounds shown in Table V were prepared by the same procedure as described above.

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EXAMPLE 12

(2S)-1-[[(2S)-2-Bis(ethoxycarbonylmethyl)amino]propanoyl]
2-pyrrolidinecarboxylic acid benzyl ester (compound 88)

Ethyl bromoacetate (0.92g) was added dropwise under ice-cooling to a stirred solution of L-alanyl-L-proline benzyl ester p-toluenesulfonate (2.24g) and triethylamine (1.53ml) in dry methylenechloride. After the addition, the reaction mixture was stirred for 2 hours at room temperature, refluxed for another 5 hours, and washed with water and saturated sodium chloride solution. The organic layer was dried over anhydrous magnesium sulfate and concentrate in vacuo. The residual oil was purified by silica gel column chromatography to give 1.02g (44.3%) of the titled

1 compound: [α]²⁴_D -67.9° (c=1.2, MeOH). IR (neat, cm⁻¹):
3460, 1742, 1642, 1428, 1180. NMR (CDCl₃, δ): 1.23 (6H, t,
J=7Hz, -CO₂CH₂CH₃), 1.25 (3H, d, J=7.2Hz, CO-CH-N), 1.67CH₃

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5 2.40 (4H, m, C_3 -H and C_4 -H), 3.57 (4H, s, -N- CH_2 CO₂Et), 3.50-4.00 (2H, m, C_5 -H), 4.13 (4H, q, J=7Hz, $-COCH_2$ CH₃), 4.10-4.67 (2H, m, C_2 -H and -CO-CH-N), 5.03, 5.20 (2H, AB_q , CH_3

10 J=12Hz, $-CH_2$ -Ph), 7.30 (5H, s, $-C_6\frac{H_5}{5}$).

The compounds shown in Table V were prepared by the same procedure as described above.

15 EXAMPLE 13

(2S)-l-[[(2S)-Bis(ethoxycarbonylmethyl)amino]propanoyl]2-pyrrolidinecarboxylic acid (compound 86)

(25)-1-[[(25)-2-bis(ethoxycarbonylmethyl)amino]propanoyl]2-pyrrolidinecarboxylic acid benzyl ester (compound 88)
(0.50g) was dissolved in ethanol (10ml), and hydrogenated
with 10% palladium on charcoal catalyst (50mg). The
titled compound was obtained as a colorless oil. Yield
0.40g (quant. yild); [a]²⁴ -52.2° (c=1.1, MeOH). IR
(neat, cm⁻¹): 1742, 1640, 1442, 1190, 1130, 752. NMR
(CDCl₃, δ): 1.23 (6H, t, J=7Hz, -CO₂CH₂CH₃, 1.25 (3H, d,
J=7.2Hz, COCH-N), 1.67-2.50 (4H, m, C₃-H and C₄-H),

1 3,53 (4H, s, N-CH₂-CO₂Et), 3.50-4.00 (2H, m, C₅-H), 4.10 (4H, q, J=7Hz, -CO₂CH₂CH₃), 4.10-4.33 (1H, m, -COCH-N), CH₃

4.47 (lH, dd, J=6.5, 5.0Hz, C_2 -H), 9.20 (lH, br. s, $-CO_2$ H).

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The compounds shown in Table V were prepared by the same procedure as described above. The following compounds are also prepared by the same procedure as EXAMPLE 12 and 13.

(2S)-1-[[4-(1-carboxy-3-pheny]propyl)amino]benzoyl]-2
pyrrolidinecarboxylic acid.

(4R)-3-[[4-(1-carboxy-3-phenylpropyl)amino]benzoyl]-2-:
(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid

EXAMPLE 14

(2S)-1-[[(2S)-2-(N-Ethoxycarbonylmethyl-N-phenylacetyl)amino]propanoyl]-2-pyrrolidinecarboxylic acid benzyl ester
(compound 90)

Phenylacetyl chloride (0.44ml) was added dropwise at

room temperature to a stirred solution of (2S)-1-[[(2S)2-(ethoxycarbonylmethyl)amino]propanoyl]-2-pyrrolidinecarboxylic
acid benzyl ester (1.1g) and triethylamine (0.47ml) in
dry acetone (15ml). After the addition, the reaction
mixture was stirred for 1 hour at the same temperature,

and the precipitate was removed by filtration. The
filtrate was evaporated in vacuo, and the residual oil was
dissolved in ethyl acetate, and washed with water and



saturated sodium chloride solution. The organic layer
was dried over anhydrous magnesium sulfate, and evaporated
in vacuo. The residual oil was purified by silica gel

column chromatography to give 1.3g (89%) of the titled compound: mp 110-110.5°C (benzene-hexane); $[\alpha]_D^{24}$ -114.0° (c=1.0, MeOH). IR (KBr, cm⁻¹): 3460, 1739, 1635, 1436, 1200, 1166. NMR (CDCl₃, δ): 1.23 (3H, d, J=7Hz, -CO-CH-N

1.28 (3H, t, J=7Hz, -CO₂CH₂CH₃), 1.67-2.50 (4H, m, C₃-H and C₄-H), 3.60 (2H, s, -COCH₂Ph), 3.33-3.90 (2H, m, C₅-H), 4.16 (2H, q, J=7Hz, -COCH₂CH₃), 4.23 (2H, s, -N-CH₂CO₂Et), 4.30-4.60 (1H, m, C₂-H), 5.03, 5.23 (2H, AB_q, J=12.5Hz, -CO₂CH₂Ph), 5.58 (1H, q, J=7Hz, -COCH-N), 7.23 (5H, s, CH₃

15 $-\text{COCH}_2\text{C}_6\text{H}_5$, 7.30 (5H, s, $-\text{CO}_2\text{CH}_2\text{C}_6\text{H}_5$).

The compounds shown in Table V were prepared by the same procedure as described above.

20 EXAMPLE 15

(2S)-1-[(2S)-?-[(1-Carboxy-3-phenylpropyl) thio]propanoyl]-2-pyrrolidinecarboxylic acid (compound 79)

(2S)-1-[(2S)-2-Mercaptopropanoyl]-2-pyrrolidine
carboxylic acid (2.0g), potassium carbonate (2.3g) and 2
bromo-4-phenylbutancic acid (2.9g) were dissolved in water

(40ml), and stirred overnight at room temperature. The

reaction mixture was acidified with 6N hydrochloric acid, and extracted with ethyl acetate. The organic layer was washed with saturated sodium chloride solution, dried over anhydrous magnesium sulfate, and concentrated in vacuo.

The residual oil was purified by silica gel column chromatography to give 2.3g (62%) of the titled compound: [\alpha]_D^{23} -82.2° (c=1.2, MeOH). IR (KBr, cm⁻¹): 1/40, 1/20, 1610, 1455, 1438, 1185, 748, 700.

The compounds shown in Table IV were prepared by the same procedure as described above.

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EXAMPLE 16

1-[[(1-Carboxy-3-phenylpropyl)amino]acetyl]-2-(2-hydroxy-phenyl)-5-pyrrolidinecarboxylic acid (compound 99)

l-(Chloroacetyl)-5-(2-hydroxyphenyl)-2-pyrrolidine-carboxylic acid [mp 204-206°C(dec.), [a] 24 +24.5° (c=1.2, MeOH)] (2.8g) was added to a stirred solution of 2-amino-4-phenylbutanoic acid (1.8g) in N sodium hydroxide (40ml). The reaction mixture was stirred overnight at room temperature. The solution was adjusted to pH 1.5 by 20% hydrochloric acid, and washed with ethyl acetate. The aqueous layer was adjusted to pH 3.0, and the separated solid was collected by filtration to give 1.0g (24%) of the titled compound. IR (nujol, cm⁻¹): 3425, 1735, 1625, 1588.



The compounds shown in Table V were prepared by the same procedure as described above.

In EXAMPLES and TABLES I, II, III, IV and V, "a" and "b" of compound No. represent diastereoisomers each other.

TABLES I, II, III, IV and V show various compounds and their physical constants including the compounds specified in EXAMPLES.

Table I

Table-continued

+		r	~			rethod of	7.07	(O.) di	(2)		IR	spectrum		Rf
No.	•	1,	T.	E	۵	prepn. (Examp. No.)	(v)	(Recrys tn. solvent)	(c, solv., °C)	Sampling*1 method	ng*1	cm_1		value (S10 ₂)
9	2-OH	ĕ	HO	0	. 2	1 5	75	190-191 (dec.) (EtOAc-MeOH)	+181.6 (1.0, McOH, 27)	£	3210, 1720, 940, 763	, 1618, 1602,	2, 1245, 1173,	0.35
7	2-OH	₹	OMe	0	7	9	83	165-166 (dec.) (EtOAc)	+164.5 (1.0, MeOii, 25)	<	3370, 1750, 755	1750, 1693, 1635,	5, 1215, 1165,	0.47
89 8	2-04	OEt	동	0	~	so	ል ·	181-182 (Etoac)	-2.8 (0.5, MeOH, 21)	. «	3310, 1727, 1190, 745	, 1703, 1637,	7, 1595, 1235,	0.55
9	2~0H	OEt	₹	0	7	s	23	116-118 (Etohc)	-311.6 (0.5, McOH, 21)	4	3370, 1735, 1180, 760	1708, 1635,	5, 1597, 1220,	0.55
9 a	2-011	ē	NIOH	0	7	7		172-173 (dec.) (EtOII-H2O)		⋖	3375, 3290, 1240, 1088,	1720, 1657, 748	7, 1625, 1590,	0.22
q 6	2-0H	₹	NHOH	0	7	7		amorph.		4	3220, 1717, 1092, 752	1655, 1625,	5, 1595, 1225,	0.33
10a	2-ਯ	OEt	NIOH	0	~	Œ		amo rph.		4	3220, 1727, 753	1625, 1595,	5, 1200,1092,	0.254
10b	2-011	OEt	NHOHN	0	7	89		amorph.						0.324
11 *5	2-OH	ē	OMe	-	~	9		amorph.	+55.5 (0.8, MeOH, 24)	ø	1738, 1630,	1585, 1310,	, 1258, 750	
11a	2-ОН	ō	OMe	7	~	9		205-207 (dec.) (benzene)	+94.6 (0.5, MeOH, 23)	æ	3110, 1730, 758	1625, 1610,	, 1192, 1121,	
12a	2-OH	Ю	픙	-	7	7	79	168-170 (dec.) (acetone-	+168.0 (0.4, MeOH, 23)	ď	3370, 1718,	1625, 1598,	, 758	0.254
12b	2-OH	퓽	₹	-	~	7		163-164 (dec.) (acetone- cyclohexane)	+149.2 (0.4, MeOH, 23)_	4	3300, 1720, 753	1708, 1615	1720, 1708, 1615, 1598, 1242,	0.254
13	2-011	ਰ	ë	0	e	ស	59	161-162 (dec.) (H ₂ O)	+153.8 (0.5, MeOil, 24)	A	3190, 1713, 943, 760	1632, 1598,	, 1253, 1098,	0.38
14	2-011	≅	OEt .	•	e e	9	98	157-158 (dec.) (ELOAc-benzene)	+145.6 (1.0, MeOH, 25)	æ	3340, 1725, 768	1638, 1597,	, 1218, 1120,	0.48
15	Ξ	ĕ	ē	0	E.	S	73	139-140 (Etuac-meoh)	+106.3 (1.0, McOil, 24)	a .	3170, 1753, 729	1709, 1631	3170, 1753, 1709, 1631, 1423, 1177,	0.39

Table-continued

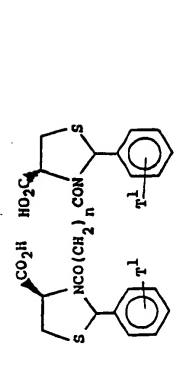
		Rf +2	value (S10 ₂)	0.31	0.43	0.47	0.50	0.52	0.55	0.56	0.57	0.57	0.57	0.51	0.57
•			-	1412,	1595,	, 1595, 850, 760		1600, (950, 760	1600, 1090,	1405,	1533, 728	1587,	1590,		1526,
		trum	cm-1	1665,	1622,	1620, 895, (1620,	1620, 1173,	1525, 735	1620, 1190,	1625, 756	1610, 775		1580, 1526, 745
		IR spectrum	ช	1710,	1700, 723	1710, 1095,		1710,	1705, 1235, 760	1615, 1095,	1663, 1240,	1660, 1043,	1655, 1042,		1660, 1050,
		Ħ	g*1	2225, 1258	3300,	3300,		3220, 1415,	3220, 1415, 830, 7	1710, 1350,	1735, 1352,	1730, 1228,	1730,		1720, 1240,
			Sampling'	æ	Ω	ø,		s a	pa	ບ່	U	ပ	٠٥		4
		.5	ĵ.	24)	24)	25)	24)	2.7)	26)	27)	23)	23)	23)	23)	23)
	Table-continued	(α)n deg.	(c, solv.,	+137.7 (1.0, MeOH,	+115.6 (1.0, MeCH,	+128.6 (0.5, MeOH,	+80.5 (1.0, MgOH,	+134.1 (0.5, NeOH,	+70.9 (0.5, MeOH,	+72.1 (0.4, MeOH,	+72.8 (1.0, MeOH,	+69.9 (0.5, MeOH,	+63.4 (0.5, MeOH,	+57.9 (0.8, Mach.,	+108.3 (0.5, MeOH,
	Table		<u>.</u>			(dec.)		(dec.)	(dec.)		•		_	_	
		(O.) du	(Recrystn. solvent)	190-191 (Etorc-Meoh)	amo rph.	158-159 (EtoAc)	oil	155-157 ((EtOAc)	153-154 ((EtOAc)	110	110	oil	tio .	oi1	amorph.
		Yield	3	29	. 62	09	33	19	63	45	79		20		45
	-		Prepn. (Examp. No.)	VA-	-	-	~	-4	~	٦	v a	~	À	٦	-
7			c	æ	4	'n	9	9	7	7	7	7	2	2	7
	}		2	0	0	0	0	•	•	0	0	0	0	0	0
		E		Ö	₹.	픙	푱	₹	ਲ	ਰ	oet	8	HO HO	푱	₹
		71	+	₹	ਰ	9	퐁	₹	ਲ	픙	픙	₹	8	품	8
		7		4-CN	2-0H	2-OH	I	2-OH	2-OH	3-NO ₂	3-NO ₂	2-F	3-E	4 - F	2-C1 5-NO ₂
		Compd	No.	16	11	18	19	20	21	22	23	24	25	56	27

Table-continued

.

m Rf a2		1710, 1620, 1600, 1410, 0.58 1230, 1090, 850, 760	1620, 1595, 0.61 1090, 943,	0.564	20, 1575, 0.52	0.534
IR spectrum	Sampling*1 method	C 1710, 1620, 1600, 14 1230, 1090, 850, 760	B 3320, 1705, 1620, 1595, 1410, 1233, 1090, 943, 850, 760		B 3280, 1700, 1620, 1575, 760,722	
- [5]	(c,	+100.3 (1.0, MGOH, 24)	slo- +120.4 (0.5, МеОН, 25)	+56.4 (0.3, MeOH, 23)	+101.4 (1.0, MeOH, 24)	+61.7 (0.6, MeOH, 23)
mp (°C)	(Recrystn.	011	123-124 (EtOAc-cyclo- hexane) (0	oi1	amorph.	of1
\$ [6] \$	(3)	68	55	26	59	43
Method of	prepn. (Examp. No.)	.		-4	7	7
	c	89	70	70	12	12
	6	0	0	0	0	0
	Ŀ	₹	₹	₹	중	ē
^	1	₹	₹	3	₹	푱
	.	2-0H	2-OH	3-CN	2-0H	3-CN
Compd t	No.	28	29	30	31	32

a and b represent diastereoisomers of the compound. A: KBr disk, B; nujol mull, C; neat. EtoAc-CHCl -AcOH (10:5:3). CHCl -EtOH-AcOH (10:2:1). EtoAc-CHCl -AcOH (7:5:1). Dicyclohexylamine salt.



Compound No. 33-37, 39-62

Compound No. 38

Compd	-		Method of	Yield	(O.) qa	[α]ν deα.		7	IR spectrum	trum			Rf +2
· 0	F	c	prepn. (Examp. No.)	3	(Recrystn. solvent)	(c, solv., °C)	Sampling*1 method	.ng*1		cm-1			value (SiO ₂)
33	2-0H	4	7	73	124-128 (MeOH)	+182.2 (1.0, DMF, 24)	es:	3280,	3280, 1726, 1620, 1596, 775,	1620,	1596,	775,	0.23
34	2-0H	S	~	67	ofi	+106.1 (0.5, MaOH, 26)	ပ	1725,	1625, 1045,	1600,	1410, 765	1235,	0.27
35	3-NO ₂	ın	4	69	111-113' (dec.) (H ₂ 0)	+88.2 (0.5, MeOH, 25)	Ø	1635, 730	1635, 1585, 1520, 1355 730	1520,	1355	1095,	0.28
36	3-CN	ĸ	м	. 65	105-112 (H ₂ 0)	+115.0 (1.0, MeOH, 25)	a ,	2270, 790	2270, 1735, 1640, 1610, 1195, 790	1640,	1610,	1195,	0.33
37	4-CN	S	М	52	amorph.	+148.2 (0.9, MeOH, 25)	Ø	2255,	2255, 1731, 1655, 1620, 785	1655,	1620,	785	0.32
36		9	7	1.1	oil	-124.5 (0.5, MeOH, 26)	ပ	1720, 880	1720, 1580, 1410, 1180, 1015, 880	1410,	1180,	1015,	0.09
39	x	9	. ~	79	amorph.	+97.4 (1.0, MeOH, 24)	ø	1720,	1720, 1625, 1585, 732	.1585,	732		0.42

Table-continued

Rf •2	(SiO ₂)	0.34	0.34	0.38	0.36	0.40	0.38	0.57	0.41	0.48	0.41	0.50	0.50	0.393
		1090,	1345,	1200,	790	1230,	1190,	1198,	1350,	1345,	790			
	_	1230,	1605, 1520,	1615,	1618,	, 1410, 763	1345,			1510,	1610,	758	767	
spectrum	см-1	1600,		1730, 1640, 1615,	1729, 1650,	1600 855,	1515, 1345,	1660, 1530, 1350,	1520, 1445,	1600, 735	1640,	1173,	1142,	
IR spe		1620, 765	1650, 730				1655,		1615, 730	1650, 1110,	1729,	1225,	1238,	•
	g*1	1720, 855,	1730, 1095,	2250, 790	2248,	1720, 1173,	1735, 730	1740, 725	1725, 1095,	1730, 1185,	2250,	1580,	1590,	•
	Sampling*1 . method	eg.	æ	a	s	ø	<u>.</u>	⋖	œ	ø.	ø	ď	≪	
	່ວ.	, 27)	21)	25)	25)	26)	25)	23)	27)	25)	25)	24)	25)	23)
(α) _n deg.	(c, solv.,	+123.6 (0.5, MeOH,	+97.5 (0.5, MeOH,	+98.3 (0.9, McOH,	+130.2 (0.9, MeOH,	+142.7 (0.5, MeOH,	+191.2 (0,6, MeOH,	+79.4 (1.0, MeOH,	+96.2 (0.5, MeOH,	+118.5 (0.5, MeOH,	+112.1 (1.1, MeOH,	+117.5 (1.0, MeOH,	+103.9 (0.5, MeOH,	+75.8 (1.0, McOH,
(D.) du	(Recrystn. solvent)	amo rph.	amorph.	amorph.	amorph.	amorph.	amorph.	61-63 (benzene)	amorph.	amroph.	amorph.	140-220 (dec.) (H ₂ O)	195-210 (dec.) (II ₂ O)	011
Yield	3	98	26	28	41	75	47	96	82	53	65	92	88	9/
Mathod	prepn. (Examp. No.)	7	~	~	8	8	8	σ	7	~	e	₹.	4	7
	c	9	ø	φ	9	7	7	7	7	7	7	7	7	►.
7	H	2-он	3-NO2	3-CN	4-CN	2-0H	2-NO ₂	3-NO ₂	3-NO ₂	4-NO	3-CN	2-F	3-F	4 - F
Compd.	NO.	40	41	42	43	4	45	46 6	47	8	49	505	512	52

Table-continued

7

Compd.	T.	c	Method of . prepn.	Pleix	mp (°C) (Recrystn.	[α] _D deg.			IR spectrum	ctrum			RE 13
į		į	(Examp.	3	Golvent)	(a, solv., *C)	Sampling method .	ng*1	:	cm-1			value (S10 ₂)
53	2-c1 5-No ₂	,	7	62	anorph.	+167.9	<	1725,		1575,	1640, 1575, 1520, 1342,	1342,	0.51
5 .	2-CH 5-SO ₂ NH ₂	~	N	. 75	amorph.		æ	1725,	/40 1620,	1595,	/40 1620, 1595, 1310, 1150,	1150,	0.424
55	2-OH	0	8	68	amorph.		æ	3300, 725	1730,	1628,	3300, 1730, 1628, 1575, 767, 725	,191	0.45
26	3-CN	©	7	41	amorph.	+104.6 (1.0, NeOH, 25)	ø	2245,	1726,	1726, 1630, 1610,	1610,	790	0.37
57	3-NO ₂	©	n	2	amorph.	+102.2 (0.5, NecH, 25)	«	1735,	1735, 1620, 1523, 1190,	1523,		728	0.47
285	3-NO2	œ	4	74	amorph.	+93.9 (0.5, MgCH, 23)	<	1597,	1520,	1269,	1597, 1520, 1269, 1096, 723	723	
59	2-0H	07	7	19	99-100.5 (dec.) (EtOAc-benzene)	+124.7 (0.5, MeCH, 27)	æ	3300,	1740,	1620,	1600, 1565, 895, 770	1565,	0.49
, 09	3-CN	10	4	63	190-195 (H ₂ 0)	+109.3 (0.5, H ₂ 0, 23)	æ	3400, 22. 778, 720	2240,		1600, 1208,	208,	
61 8 8	2-0ii	75	71	99	amorph.	+69.5 (1.0, MeCH, 24)	Ø	3300, 725	3300, 1728, 1630, 725	1630,	1590, 762,	62,	0.45
· 62	3-CN	12	4	25	amorph.	. +104.2 (0.5, Mach, 23)	Ø	3400, 222: 775, 720	2225, 1	1605, 1	3400, 2225, 1605, 1320, 1207, 775, 720	207,	0.463

1 A; KBr disk, B; nujol mull, C; neat.
2 EtoAc-CHCl3-AcOH (10:5:3),
3 EtoAc-CHCl3-AcOH (7:5:1),
4 CHCl3-McOH-AcOH (3:1:1).
5 Disodium salt,
5 Dimethyl ester.

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со2и но₂с

Compound No. 69-71

Compound No. 63-68

Compd.			Method of	Yield	() du	(α) n Jea.		IR	IR spectrum	rcm		Rf #2
		3	prepn. (Examp. No.)	3	(Recrystn. solvent)	(c, golv., °C)	Sampling*1 method	1941	cm-1	7		value (SiO ₂)
]]	2-ОН	-сн ₂ сосн (сосн ₃) -	v	31	amorph.		8	1743,	l .	1720, 1630, 1600,	1600,	0.383
) - ;	2-OH	-CH ₂ -0-CH ₂ -	7	35	amorph.	(1.2, MeOH, 25) +138.6 (1.1, MeOH, 25)	«	3300,	1726,	3300, 1726, 1640, 1453, 1234, 1142	1453,	0.24
-	NO ₂	3-NO2 +CH25 (O) +CH2+2	-	36	amorph.	+81.7 (0.9, MaCH, 24)	<	3400,	1702,	1618, 1525,	1525,	0.553
) - 2	2-0iI	1CH2120-1CH212-	7	33	136-137 (Etoac)	+147.6 (0.5, MaOH, 25)	æ	3320, 1595, 770	1750, 1235,	1710, 1110,	1625, 855,	0.28
) - 2	2-0il	tcH ₂ 12-(CH ₂ 12-	-	40	159-160 (dec.) (EtOAc)	+136.4 (0.5, HeOH, 27)	æ	3360, 1435, 763	1710, 1235,	3360, 1710, 1627, 1599, 1435, 1235, 1099, 852, 763	1599, 852,	0.43
2-C	2-OH	(CH ₂) 5 - (CH ₂) 5 - (CH ₂) 7	-	35	amorph.	+78.1 (1.0, NeOH, 24)	. · ·	3300, 760	1715,	3300, 1715, 1627, 1590, 760	1590,	0.31
Z	NO ₂	3-NO ₂ +CH ₂ +2(O)+CH ₂ +2	7	. 44	amorph,	+106.9 (1.1, MeOH, 24)	• • .	3425,	1730, 1350	3425, 1730, 1640, 1525, 1400, 1350	1525,	0.38

Table Il

Table-continued

			Mathod	# F 0 7 %	(D.) de	מפט ענשן		IR spectrum	Rf *2 . value
Compd. T.	- [+	3	prepn. (Examp. No.)	3	(Recrystn. solvent)	(c, solv., •C)	-Sampling*1 method	cm-1	(8102)
70	. 2-OH	70 2-ОН 4CH ₂ 1 ₂ O-4CH ₂ 1 ₂	7	47	47 amorph.	+83.0 - (0.5, MeOH, 26)	B 1720 850,	1720, 1625, 1600, 1230, 1090, 850, 760	0.15
n	2-OH	2-on tcH2125+CH212	8	. 53	amorph.	+129.3 (0.5, MeCH, 27)	B 1720 1093	1720, 1620, 1600, 1420, 1230, 1093, 852, 763	0.30

*1 A; KBr disk, B; nujol mull. *2 EtOAc-CHCl₃-AcGf (10:5:3). *3 EtOAc-EtOH-AcOH (40:1:1). *4 CHCl₃-EtOH-AcOH (10:2:1).

Compound No. 77-80

Compound No. 72-76

1		1					•	
Rf •2	value (S10 ₂)	0.26	0.223	0.74	0.65	0.134	0.134	99.0
		3030, 1737, 1720, 1615, 1413, 1215, 1150, 717	1735, 1623, 1413, 1243, 1170, 1043, 699	1400,	700	756	755	1735, 1630, 1615, 1420, 1242,
		1615,	1243,	1600,	750,	1217,	1215,	1420,
IR apectrum	cm-1	1720,	1413,	1720-1710, 1625, 1600, 1400, 1235, 752, 698	1720, 1620, 1415, 750, 700	1720, 1620, 1422, 1217, 756	1722, 1620, 1420, 1215, 755	1615,
IR SP		1737,	1623, 699	1710, 1 752, 6	1620,	1620,	1620,	1735, 1630, 161
	ing*1	3030,	1735,	1720-1710, 1235, 752,	1720,	1720,	1722,	1735,
	Sampling*1 method	<	υ	<	<	ပ	U	ပ
5	ΰ.	23)	23)	25)	25)	25)	25)	241
[α] _D deg.	(c, solv., °C)	+8.6 (1.0, MeOH, 23)	-161.5 (1.0, MeOH,	+122.1 (1.2, HeOH,	-97.9 (1.1, MaCH,	-52.2 (1.2, MeOH, 25)	-60.4 (1.0, MaOH,	-61.2 (1.3. MeOH. 24)
Yield mp (°C)	solvent)	151-153 (EtoAc)	041	amorph.	amorph. (011	110	011
Yield	દ	38	49	18	25	37	46	84
Method of prepn.	(Examp. No.)	10	10	10	.10	70	10	10
LT		z	z	Ξ	I	x	x	×
ه د		4 4	Ph Ph	CII ₂ CH ₂ Ph	CH ₂ CH ₂ Ph	x	I	x
بار م		CH ₃	CH ³	×	ਜੂ ਜੂ	CH ₂ Ph	CH ₂ Ph	CH ₂ CH ₂ Ph
₹.		×	≖ ≺		I	I	I	I
Compd.	<u>.</u>	72a	72b) ودر	74	75a	75b	. 92

= :	07 5	8	151-153 (BtOAc)	+8.6 (1.0, MgOH, 23)	4	3030, 1737, 1720, 1615, 1413, 1215, 1150, 717	, 1720,	1615,	1413,	0.26
	0	4	041	-161.5 (1.0, MeOH, 23)	o a	1735, 1623, 1413, 1243, 1170, 1043, 699	, 1413,	1243,	1170,	0.22
	01	81	amorph.	+122.1 (1.2, MeOH, 25)	٠	1720-1710, 1625, 1600, 1400, 1235, 752, 698	1625, 698	1600,	1400,	0.74
	.10	25	amorph.	-97.9 (1.1, MaCH, 25)	۷ -	1720, 1620, 1415, 750, 700	, 1415,	750,	007	0.65
	10	37	oil	-52.2 (1.2, MeOH, 25)	υ •	1720, 1620, 1422, 1217, 756	, 1422,	1217,	756	0.134
	10	46	011	-60.4 (1.0, MaOH, 25)	υ •	1722, 1620, 1420, 1215, 755	, 1420,	1215,	755	0.134
	10	84	011	-61.2 (1.3, MeOH, 24)	υ.	1735, 1630, 1615, 1420, 1242, 1172, 1043, 702	, 1615, , 702.	1420,	1242,	99.0

Table-continued

+ + +	v	ve	_	Method of	Yield	mp (C°)				IR spectrum	rum		Rf 12
No.	÷). L	È-	prepn. (Examp. No.)	3	(Recrystn. solvent)	(c, Bolv., °C)	ត	Sampling*1 method		cm-1		value (S10 ₂)
. 11	æ	COPh	瓦比	15	36	o11	-46.2		c 1733,	1733, 1678, 1632, 1610, 1447,	32, 161	0, 1447,	0.323
78	×	CH2CH2Ph	±	15	46	011	(0.8, MeOH, 30) -48.4 (1.1, MeOH, 26)	(9) (9)	1258, 118 C 1730, 161 750, 703	1258, 1187, 1025, 1001, 751 1730, 1610, 1450, 1240, 1190, 750, 703	25, 100 50, 124	1, 751 0, 1190,	0.72
62	<u>5</u>	CH ₃ CH ₂ CH ₂ Ph	×	15	62	amorph.	-82.2 (1.2, MeOH, 23)	(2)	A 1740,	1740, 1720, 1610, 1455, 1438, 1185, 748, 700	10, 145	5, 1438,	0.38
. 08	×	E HOOO	Et.	15	45	: 110	-49.6 (0.9, MeOH, 30),	, (0,	c 1736,	1736, 1597, 1398, 1378, 1333, 1250, 1191, 1047, 860, 752	98, 137 47, 860	8, 1333, , 752	0.293

a and b represent diastereoisomers of the compound. A; KBr disk, C; neat. EtOAc-CHCl3-AcOH (10:5:3). Benzene-EtOAc-EtOH-AcOH (14:14:2:1). Benzene-EtOAc-AcOH (25:25:1). CHCl3-EtOH-AcOH (10:2:1)

Table V

.

Compound No. 99

Compound No. 86-98, 100-102

Compound No. 81-85

RE	(S10 ₂)	0.252	0.45	0.744	0.694		0.215
,				-		16 14 00	
· whi	cm-1	3200, 1672, 1440, 1335, 752	3420, 3210, 1650, 1240, 839, 790	3370-2900, 1655, 1602, 1175	1720, 1644, 744	1720, 1660, 1492, 1452, 752, 700	-
IR spectrum		3400, 1740, 1560, 1380,	3420, 32 1650, 12 839, 790	3370- 1655, 1175	3350, 1670, 1236,	3400, 1610, 1240,	1742, 1442, 1130,
IR	Sampling method	~	ø	⋖	4	4	ပ
		24)	23)	6	<u> </u>		=
ob dea	(c, solv., °C)	+271.2 (0.5, N NAOH; 24)	+94.7 (0.5, N NAOH, 23)	+86.5 MeOH, 26)	+78.9 MaCH, 25)		-52.2 MeOH, 24)
2	(c, sc	+27 (0.5, 18	6+ (0.5, N	+86.5 (0.4, MeOH,	+78.9 (0.8, MaCH,		-52.2 (1.1, MeOH,
	• .	lec.)		lec.) ir)	lec.)	ec.)	:
()•) dist	(Recrystn. solvent)	181-182 (dec.) (H ₂ O)	150-155 (H ₂ 0)	150-153 (dec.) (BtOH-ether)	50.3: 172-173 (dec.) (EtOAc)	174-175 (dec.) (H ₂ 0)	#
8	(Re	181-18 (H ₂ O)	150-1 (H ₂ 0)	150 (Bt	172. (Et	174-1 (H ₂ 0)	011
Yeard	3	48.2	32.8	44.8	50.3	27.2	quant.
Method	prepn. (Examp. No.)	п	a	n	11	11	13
O.	£ 4	±.	z.	35	z	=	×
				1 '		CH ₂ CH ₂ Ph -CH-	•
ď	, E	-сн ₂ -	-G			-CH-	-CH2-
							Cil ₂ CO ₂ Et
4	£-	×	I	x	I	=	
	F	=	×	x	x =	=	Et
4	£	* -	⊘	₹	₹ ⊙	₹ -	z
+ pomod	NO.	81	85	93	84	88	98 .

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.

Table-continued

Compd.t No.	4 t	T T8	o. F•	T.10	of of prepn. (Examp.	Xield (*)	mp (°C) (Recrystn. solvent)	[α] _D deg. (c, solv., °C)	Samp	IR spectrum ling cm ⁻¹	Rf value (SiO ₂)
87	= =	CH ₂ CO ₂ H	-CH ₂ -	x	7	26	amorph.	-32.8 (1.0, MeOil, 24)	øs.	3400, 1720, 1640, 1460, 1380	0.102
88	H	Et CH ₂ CO ₂ Et	후 (CH ₂ Ph	12	45.2	oil .	-67.9 (1.2, MeOH, 24)	υ		0.70
89a . 1	H H	I	Q (×	16	33	216-218 (dec.) (H ₂ 0)	-141.1 (0.3, MeCH, 23)	Ø	2600, 1743, 1550, 1250, 1230, 800	0.20
ਜ વ 68	I I	=		r	16	45	218-226 (dec.) (H ₂ 0)	+1.5 (0.5, MeOil, 23)	æ		0.200
06	H	t COCH ₂ Ph	-G ₂ -	CH ₂ Ph	7.	68	110-110.5 -114.0 (benzene-n-hexane) (1.0, MeOH,	-114.0 B) (1.0, MeOH, 24)	«	3460, 1739, 1635, 1436, 1200, 1166	0.457
	H Et	t COCH ₂ Ph	-CH ₂ -	r	13	guant.	. oll	-99.7 (1.1, MeOH, 23)	۵		0.355
92	I I	COCH ₂ Ph	-CH ₂ -	x	7	83	205-206 (Etoac-Meoh)	-123.5 (1.0, MeOH, 24)	4	3430, 1727, 1635, 1598, 1426, 1184	0.388
н [6	E E			CH ₂ Ph	14	93	011	-93.2 (1.0, MeOH, 24)	U .	1655, 1188	0.517
				I	13	quant.	ott	-94.7 (1.2, MeOH, 23)	Ω	1746, 1642, 1449, 1190,	0.38
95 #	=	CO(CH ₂) ₂ Ph	-CH ₂ -	=	1	96	amorph.	-104.3 (1.0, MeOH, 24)	4	3440, 1735, 1610, 1450, 1185	0.458
96	l Et	: CH ₂ Ph	-CH ₂ -	CH ₂ Ph	12	46	oil	-66.0 - (1.2, Madi, 25)	۵	1740, 1639, 1450, 1425, 1185	0.57
		CH ₂ Ph	-CH ₂ -	×	7	87	amorph.	-59.0 (1.1, MeOH, 25)	4	3420, 1720, 1638, 1448, 1385	0.172
86 H	=		CII ₂ CII ₂ Ph CHCO ₂ H	x	14	62	195-196 (dec.) (EtOAc)	•	.	1758, 1720, 1615, 1600, 1380, 750,	

Table-continued

Ş

Compd. T4	7	е _н	O _E	10	of of prepn.	Yield	mp (°C) (Recrystn.	(a) G deg.	H	IR spectrum	3
					(Examp.	3	solvent)	(c, solv., °C)	Sampling method	ng cm-1	(S10 ₂)
11,66	¥ 80	CH ₂ CH ₂ Ph CHCO ₂ H	31	×	16	24	amorph.		æ	3425, 1735, 1625, 1588	0.662
н .	n n	T	. (*)	CH ₂ Ph	7.	37	011	-46.9 (0.5, MeOH, 23)	U	1740, 1642, 1453, 1425, 1170, 740	0.20
н 101	gi Ti	8 N N N N N N N N N N N N N N N N N N N		±	13	06	ofl	-35.9 (0.5, MeOH, 23)			0.25
102 н	I	2 6 K	-2 -2 -2	æ	,	06	228-230 (dec.) (MeOH)	-33.9 (0.4, MeOH, 23)	æ	3450, 1720, 1610, 1305, 1228, 1200, 680	0.3410

and b represent diastereoisomers of the compound.

mp 204-206°C (dec.), [α]²⁴ +24.5° A: KBr disk, B: nujol mull, C: Neat, D: liquid cell (CHCl₃).

n-BuOH-AcOH-H₂O (4:2:1).

n-BuOH-AcOH-H₂O (4:1:2).

EtOAc-CHCl₃-AcOH (10:5:3).

EtOAc-CHCl₃-AcOH (7:5:1).

EtOAc-CHCl₃-AcOH (7:5:1).

Denzene-EtOAc-AcOH (25:25:1).

CHCl₃-EtOH-AcOH (10:2:1).

EtOAc-CHCl₃-AcOH (10:2:1).

CHCl₃-EtOH-AcOH (10:2:1).

Starting material: 1-(chloroacetyl)-5-(2-hydroxyphenyl)-2-pyrrolidinecarboxylic acid; (c=1.2, MeOH), IR (nujol, cm⁻¹) 3370, 1698, 1645, 1610, 1595, 1238, 758.

1 PHARMACOLOGICAL TEST 1

It has been known that aldose reductase participates in diabetic cataract which is one of the diabetic complications and that appearance is retarded or depressed by inhibition of the aldose reductase [Acta Societatis Ophthalmologicae Japonicae, 80, 1362 (1976)]. The following method is used for the present test.

(Method)

Aldose reductase is purified from rat lenses according to the method of Hoyman et al. [J. Biol. Chem., 240, 877 (1965)]. Action of the compounds (I) of this invention is evaluated by measurement of optical density according to the J.H. Kinoshita's method [Invest. Ophthal., 13, 713 (1974)]. The reaction mixture for the measurement of the aldose reductase activity is 3.0ml [0.007M phosphate buffer solution (pH 6.2), 0.46M lithium sulfate, 5 x 10⁻⁵M NADPH, 4 x 10⁻⁴M DL glyceraldehyde, 10U aldose reductase, 10⁻⁴ to 10⁻¹⁰M the compounds (I)] as total volume, and the absorbance thereof is measured at 340nm.

(Result)

Table VI shows that the compounds (I) of this in25 vention have a strong aldose reductase inhibition effect.

1 Table VI. Inhibitory Activity of the Thiazolidine Compounds against Aldose Reductase

5	Compd. No.	IC ₅₀ (M)*1
J	22	8.2×10^{-10}
	23	1.1×10^{-8}
	47	1.6×10^{-10}
	56	1.7×10^{-9}
10	57	5.4×10^{-9}
•	Control ^{*2}	1.0×10^{-7}

Molar concentration of a compound producing 50% inhibition of aldose reductase.

Quercitrin: referred to Acta Societatis
Ophthalmologicae Japonicae, 80, 1369-1370 (1976).

PHARMACOLOGICAL TEST 2

• ••

converting enzyme activity, bioassay for the contractile response of isolated smooth muscle or the pressor response of normal animals and biochemical assay for the enzyme isolated from lung or other organs of animals are known. The former is found more advantageous than the latter for the examination of the convertion of angiotensin I to angiotensin II in vivo.

In the present study, therefore, we adopted the bioassay for contractile response of isolated guinea pig ileum to angiotensin I.

5 (Method)

25

Isolated guinea pig ileum was suspended in the organ bath containing 20ml of Tyrode's solution of 30°C gassed with 95% O₂ + 5% CO₂. The contraction induced by the addition of angiotensin I (0.1µg/ml) at intervals of 10 minutes was recorded on a recticorder (Nihon Koden) for 90 seconds using FD pick up (ST-1T-H, Nihon Koden)

The test compounds were added to the bath 5 minutes

before the addition of angiotensin I.

The inhibitory activity of angiotensin I-converting of the enzyme was calculated by the following formula.

$$\frac{A - B}{A} \times 100$$

- A: contractile intensity of angiotensin I before addition of the compound
- B: contractile intensity of angiotensin I after addition of the compound

From the fact that kininase II, which destroys bradykinin having contractive action on isolated guinea pig ileum, is thought to be identical with angiotensin I-converting enzyme augmentation of the contractile response to bradykinin by test compounds was examined

by using bradykinin (0.005μg/ml) in place of angiotensin I according to the above mentioned method. (Result)

Concentration of a number of the compounds of this invention, which produced 50% inhibition of angiotensin I activity or augmentation of bradykinin activity inducing the contraction of guinea pig ileum, fell in the range of 10^{-7} – 10^{-9} M.

10 PHARMACOLOGICAL TEST 3

The activity of angiotensin I-converting enzyme was measured by spectrophotometry according to the method of D.W. Cushman and H.S. Cheung [Biochem. Pharmacol., 20, 1637 (1971)]. That is, the absorbance of hippuric acid was measured, which is liberated by incubating hippuryl-L-histidyl-L-leucine (HHL) as substrate in the presence of angiotensin I-converting enzyme extracted from rabbit lung.

20 (Method)

The reaction mixture is as follows:

100mM phosphate buffer (pH 8.3)

300mM sodium chloride

5mM HHL

 $10^{-3} \sim 10^{-9} \text{M enzyme inhibitor}$ 5 mU enzyme

1 0.25ml of the above mixture was incubated at 37°C for 30 minutes and the reaction was stopped by adding 0.25ml of 1 N hydrochloric acid. To this solution, 1.5ml of ethyl acetate was added in order to extract hippuric acid. 1.0ml of ethyl acetate layer was collected and evaporated to dryness, and the residue obtained was dissolved in 1.0ml of water. The absorbance of this solution was measured at 228nm.

The inhibitory activity of angiotensin I-converting 10 enzyme was calculated by the following formula:

Percent inhibition = $\frac{A - B}{A} \times 100$

A: absorbance of reaction solution before addition of the compound

B: absorbance of reaction solution after addition of the compound

Concentration of compound producing 50% inhibition of angiotensin I-converting enzyme (IC_{50})

The solution containing compounds at the concentra-20 tion of $1 \times 10^{-3} M$ to $1 \times 10^{-9} M$ was incubated and percent inhibition at each concentration was calculated according to the above formula, and then IC_{50} , concentration of the compound producing 50% inhibition of the enzyme activity, was determined.

25 (Result)

15

IC50 of a number of the compounds of this invention,

1 fell in the range of $10^{-7} \sim 10^{-10} M$.

TOXICITY TEST

The acute toxicity of compounds 47 and 56 is 1000 ~ 5 1500mg/kg.

(Experimental animals)

The male ddy-std. strain mice (4 weeks of age, weighing 19-21g) were placed in a breeding room of con
stant temperature and humidity (23+1°C, 55+5%) and fed freely pellet diet (CE-2, Clea Japan, Inc.) and water ad. libitum for a week. The mice showing the normal growth were selected for the experiment.

15 (Method of administration)

7

Test compounds are dissolved in distilled water and administered (i.v.) in a dose of 0.5ml/20g body weight.

It is found in the above pharmacological and toxicity test that the compounds (I) of this invention are useful as drugs for therapy or prophylaxis of the diabetic complications and as antihypertensive agents.

In case the compounds are used for preventing or relieving diabetic complications, the dosage forms are tablet, capsule, granule, powder, suppository, injection,

ophthalmic solution, ophthalmic ointment, etc. These proparations can also contain general excipients.

for reducing blood pressure, they can be given with the combination of diuretics such as probenecid, carinamide, hydroflumethiazide, furosemide, and bumetanide same as other antihypertensive agents. The compounds can be administered either orally or parenterally. The dosage forms are tablet, capsule, granule, powder, suppository, injection, etc. In the treatment of hypertension, these preparations can contain not only general excipients

10 but also other antihypertensive agents such as reserpine, α-methyldopa, guanethidine, clonidine, hydralazine, etc., or β-adrenergic blocking agents such as propranolol, alprenolol, pindolol, bufetolol, bupranolol, bunitrolol, practolol, oxprenolol, indenolol, timolol, bunolol, etc.

The dose is adjusted depending on symptom, dosage form, etc. But, usual daily dosage is 1 to 5000mg, preferably 10 to 1000mg, in one or a few divided doses.

EXAMPLES OF FORMULATION

20 (1) Oral drug

(a) tablet

	Total	240mg
	magnesium stearate	3mg
25	calcium carboxymethylcellulose	7mg
	crystalline cellulose	60mg
	lactose	120mg
•	compound 13	50mg

4		
1	compound 22	100mg'
	lactose	95mg
	crystalline cellulose	45mg
	calcium carboxymethylcellulose	7mg
5	magnesium stearate	3mg ·
	Total	240mg
	compound 23	150mg
10	lactose	60mg
10	crystalline cellulose	30mg
	calcium carboxymethylcellulose	7mg
	magnesium stearate	3mg
	Total	250mg
15		- · · · · · · ·
, ,	compound 56	150mg
	lactose	60mg
	crystalline cellulose	30mg
	calcium carboxymethylcellulose	7mg
20	magnesium stearate	3mg
	Total	
•		250mg
•	compound 74	150
	lactose	150mg
25	crystalline cellulose	60mg
£.J		30mg
	calcium carboxymethylcellulose	7mg

.

1	magnesium stearate	3mg,
	Total	250mg
	•	
	compound 88	150mg
5	lactose	60mg
	crystalline cellulose	30mg
	calcium carboxymethylcellulose	7mg
	magnesium stearate	3mg
10	Total	250mg

The tablets may be treated with common film-coating and further with sugar-coating.

15	(b)	granule	
15		compound 13	30mg
		polyvinylpyrrolidone	25mg
		lactose ·.	385mg
		hydromypropylcellulose	50mg
20		talc	10mg
		Total	500mg
		compound 22	30mg
		polyvinylpyrrolidone	25mg
25		lactose	385mg
		hydroxypropylcellulose	50mg

1	talc	10mg.
	Total	500mg
_	compound 94	30mg
5	polyvinylpyrrolidone	. ² 25mg
	lactose	385mg
	hydroxypropylcellulose	50mg
	talc	10mg
10	Total	500mg
	(c) powder	•
	compound 13	250mg
	lactose	240mg
15	starch	480mg
13	colloidal silica	30mg
	Total	1000mg
-	compound 65	300mg
20	lactore	230mg
	starch	440mg
•	colloidal silica	30mg
	Total	1000mg
25	compound 79 lactose	300mg 230mg

1	starch	440mg
	colloidal silica	30mg
	Total	1000mg
5	·	••
3	compound 100	300mg
	lactose	230mg
	starch	440mg
	colloidal silica	30mg
10	Total	1000mg
		•
(d) capsule	
	compound 13	50mg
	lactose	102mg
15	crystalline cellulose	36mg
	colloidal silica	2mg
	Total	190mg
	compound 23	100mg
20	lactose	52mg
	crystalline cellulose	36mg
	colloidal silica	. 2mg
	Total	190mg
25	compound 74	200mg
	glycerin	179.98mg

1	butyl p-hydroxybenzoate	0.02mg
	Total	380mg
5	compound 81	30mg
5	glycerin	349.98mg ·
	butyl p-hydroxybenzoate	0.02mg
	Total	380mg
10	compound 98	200mg
	glycerin	179.98mg
	butyl p-hydroxybenzoate	0.02mg
	Total	380mg

15 (2) Injection

- (a) 1 to 30mg of compound 9B is contained in 1ml of the aqueous solution (pH 6.5-7.0).
- (b) 1 to 30mg of compound 73 is contained in 1ml of the aqueous solution (pH 6.5-7.0).
 - (3) Ophthalmic solution

The following composition is contained in 5ml of the aqueous solution (pH 6.0).

25.

Compound 23

50mg

1		<pre>propyl p-hydroxybenzoate</pre>	0.7mg
		methyl p-hydroxybenzoate	1.3mg
		sodium hydroxide	proper quantity
5	(4)	Ophthalmic ointment	•
		The following composition	is contained in 1g.
		•	
		compound 22	20mg
		white petrolatum	889.8mg
10		mineral oil	100mg :
		butyl p-hydroxybenzoate	0.2mg
		_	
	(5)	Suppository	
		The following composition	is contained in 1g.
15			
		compound 47	50mg
		polyethylens glycol 1000	800mg
-		polyethyler = glycol 4000	150mg
			•

20

25

1. A compound of the formula [I]

wherein

 Q^1 and Q^2 each is methylene or sulfur, but Q^1 and Q^2 are not sulfur at the same time;

RA is Ra or Rb:

RB and RC each is RC;

W is
$$\begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}_{z} \begin{bmatrix} R^3 \\ C \\ R^4 \end{bmatrix}_{m} \times \begin{bmatrix} R^5 \\ C \\ R^6 \end{bmatrix}_{n} \begin{bmatrix} R^7 \\ C \\ R^8 \end{bmatrix}_{p} \times \begin{bmatrix} R^9 \\ C \\ R^{10} \end{bmatrix}_{q} \begin{bmatrix} R^{11} \\ C \\ R^{12} \end{bmatrix}_{r} \times \begin{bmatrix} R^{13} \\ C \\ R^{14} \end{bmatrix}_{s} \begin{bmatrix} R^{15} \\ R^{16} \end{bmatrix}_{t}$$
, wherei

X,Y and Z each is single bond, $-CH_2^-$, $-C = C_-$,

1, m, n, p, q, r, s and t each is 0, 1, 2 or 3;

R¹, R², R³, R⁴, R⁵, R⁶, R⁷, R⁸, R⁹, R¹⁰, R¹¹, R¹², R¹³, R¹⁴, R¹⁵, R¹⁶, R¹⁷, R¹⁸,

R¹⁹, R²⁰ and R²¹ each is R^d;

R²³

A. _b.

 R^{A} is R^{b} when W is -CH-NH-C- or -CH-(CH) wherein R^{22} , R^{23} , R^{24} , R^{25} and R^{26} each is R^{d} ;

25

P

15

20

Ra is selected from the group consisting of

- (i) hydrogen, lower alkyl and lower alkenyl, and
- (ii) lower alkyl and lower alkenyl substituted by at least one substituent selected from the group consisting of lower alkyl,

- lower alkenyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto. acylmercapto, lower alkylthio, carbox lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
- Rb is selected from the group consisting of

 (a)(i) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl, and

 (ii) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen, nitro, cyato, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl, and (iii) carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl and heteroaryloxycarbonyl;
- (b) (i) phenyl and naphthyl, and

 (ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylamino-sulfonyl and lower alkylsulfinyl;
- (c) (i) furyl, thienyl and pyridyl, and
 (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxy-carbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

1

R is selected from the group consisting of

(a) (i) hydroxy, lower alkoxy and amino, and

(ii) lower alkoxy and amino substituted by at least one substituen

selected from the group consisting of lower alkyl, aralkyl, heteroaralkyl, aralkenyl, heteroaralkenyl, hydroxy, lower alkoxy, aralkyloxy, heteroaralkyloxy, aryloxy, heteroaralyl, acyloxy, aryl, heteroaryl, substituted aralkyl and substituted aryl wherein the substituent is lower alkyl, lower alkoxy, halogen or amino;

(b)(i) aryloxy and heteroaryloxy, and

(ii) aryloxy and heteroaryloxy substituted by at least one substituent selected from the group consisting of lower alkyl, hydroxy, lower alkoxy, halogen and amino, and

CO-RB
Q2
NQ1
RA

10

R^d is selected from the group consisting of

(a) (i) hydrogen, lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo, and

(ii) lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, lower alkoxy, lower alkanoyl, aryl, heteroaryl, acyloxy, aroyl, hydroxy, carboxy, amino, guanidino, mercapto, acylamino, acylmercapto, lower alkoxycarbonyl, sulfo, halogen, nitro, cyano, sulfamoyl, lower alkylaminosulfonyl, lower alkylthio and lower alkylsulfinyl;

(b) (i) phenyl and naphthyl, and (ii) phenyl and naphthyl substituted by at least one substituent lower alkanoyl, acyloxy, hvdroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl (c)(i) furyl, thienyl and pyridyl, and

(ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylene-dioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkyl-aminosulfonyl and lower alkylsulfinyl;

and salts thereof.

5

- 2. A compound of claim 1 wherein $-Q^1-Q^2$ is $-CH_2CH_2$, $-SCH_2$ or $-CH_2S$.
- 3. A compound of claim 1 wherein R^a is hydrogen, methyl, othyl, 1-methylethyl, propyl, 2-methylpropyl, butyl, 2,6-dimethyl-5-heptenyl, cyclohexyl, S-acetyl-2-mercaptoethyl or 2-mercaptoethyl.
- 4. A compound of claim 1 wherein R^b is benzyl, 2-phenylethyl, 4-methylbenzyl, 4-methoxybenzyl, 2-hydroxybenzyl, 4hydroxybenzyl, 3-fluorobenzyl, 3-nitrobenzyl, 3-cyanobenzyl,
 2-(4-methoxyphenyl)ethyl, 2-(2-hydroxyphenyl)ethyl, 2-(4hydroxyphenyl)ethyl, 2-(3-fluorophenyl)ethyl, 2-[3-(trifluoromethyl)phenyl]ethyl, 2-(3-nitrophenyl)ethyl, 2-(3-cyanophenyl)ethyl, 2-pyridylmethyl, 4-pyridylmethyl, 2-furylmethyl, 2-(2pyridyl)ethyl, 2-(4-pyridyl)ethyl, 2-(2-furyl)ethyl, phenyl,
 4-methylphenyl, 2-chlorophenyl, 4-chlorophenyl, 2,4-dichlorophenyl,
 2-fluorophenyl, 3-fluorophenyl, 4-fluorophenyl, 2-nitrophenyl,
 3-nitrophenyl, 4-nitrophenyl, 2-chloro-5-nitrophenyl, 4-dimethyl-

- aminophenyl, 4-acetaminophenyl, 4-[(benzyloxycarbonyl)amino]chenyl,
 2-carboxyphenyl, 4-carboxyphenyl, 2-hydroxyphenyl, 3-hydroxyphenyl, 4-hydroxyphenyl, 3-benzoxyphenyl, 4-(benzyloxycarbonyloxy)phenyl, 3,4-dihydroxyphenyl, 5-chloro-2-hydroxyphenyl, 2-methoxyphenyl, 4-methoxyphenyl, 3,4-dimethoxyphenyl, 3,4,5-trimethoxyphenyl, 2-hydroxy-3-methoxyphenyl, 2-hydroxy-4-methoxyphenyl,
 4-hydroxy-3-methoxyphenyl, 3,4-methylenedioxyphenyl, 2-cyanophenyl, 3-cyanophenyl, 4-cvanophenyl, 2-nitrosophenyl, 3nitrosophenyl, 4-nitrosophenyl, 2-hydroxy-5-sulfamoylphenyl,
 2-hydroxy-5-[(diprop;lamino)sulfonyl]phenyl, 3-(methylsulfinyl)phenyl,
 3-(difluoromethoxy)phenyl, 1-naphthyl, 2-furyl, 2-(5-methyl)furyl,
 2-thienyl, 3-pyridyl or 4-pyridyl.
 - 5. A compound of claim 1 wherein R^C is hydroxy, methoxy, ethoxy, butoxy, amino, hydroxyamino, succinimidomethoxy, 1-succinimidoethoxy, phthalimidomethoxy, 2-phthalimidoethoxy, pivaloyloxymethoxy, 1-pivaloyloxyethoxy, benzyloxy, phenoxy, benzyloxyamino or 0-0 R^A N CO-R^B
- 5 A compound of claim 1 wherein R^d is hydrogen, methyl, ethyl, propyl, 1-methylethyl, 2-methylpropyl, 4-methylpentyl, vinyl, allyl, 2-butenyl, 1,3-butanedienyl, 1-methylvinyl, hydroxymethyl, carboxymethyl, 2-carboxyethyl, cyclohexyl, cyclohexylmethyl, benzyl, 2-phenylethyl, 3-phenylbutyl, 2-(1-naphthyl)ethyl, 2-(4-chlorophenyl)ethyl, 2-(3,4-dichlorophenyl)ethyl, 4-methoxybenzyl, 2-(4-methoxyphenyl)ethyl, i-hydroxybenzyl,) 2-(4-hydroxyphenyl)ethyl, (2-pyridyl)methyl, (4-pyridyl)methyl, 2-(2-pyridyl)ethyl, 2-(4-pyridyl)ethyl, (4-imidazolyl)methyl, 3-indolylmethyl, 2-(methylthio)ethyl, 4-aminobutyl, 5-aminopentyl, 4-guanidinobutyl, 4-(aminomethyl)benzyl, phenoxymethyl, (phenylthio) methyl, l-amino-2-phenylethyl, l-amino-3-methylb..tyl phenyl, naphthyl, 4-methylphenyl, 2-chlorophenyl, 4-chlorophenyl, 2,4-dichlorophenyl, 2-fluorophenyl, 3-fluorophenyl, 4-fluorophenyl, 2-nitrophenyl, 3-nitrophenyl, 4-nitrophenyl, 2-chloro-5-nitrophenyl, 4-dimothylaminophenyl, 4-acetaminophenyl, 2-carboxyphenyl, 4-carboxyphenyl, 2-hydroxyphenyl, 3-hydroxyphenyl, 4-hydroxyphenyl, 3-benzcxyphenyl, 3,4-

dihydroxyphenyl, 5-chloro-2-hydroxyphenyl, 2-methoxyphenyl,
4-methoxyphenyl, 3,4-dimethoxyphenyl, 3,4,5-trimethoxyphenyl,
2-hydroxy-3-methoxyphenyl, 2-hydroxy-5-sulfamoylphenyl, 3(methylsulfinyl)phenyl, 3-(difluoromethoxy)phenyl, 2-furyl, 2-(5-methyl)furyl, 2-thienyl, 3-pyridyl or 4-pyridyl.

5

7. A compound of claim 1 wherein W is
$$-CH - (CH_2)_{0-12} \frac{CH}{R^{16}}$$
, $-CH - (CH_2)_{0-6} \frac{CH}{R^{17}} \frac{CH}{R^{18}}$, $-CH - (CH_2)_{0-6} \frac{CH}{R^{16}}$, $-CH - (CH_2)_{0-4} \frac{CH}{R^{17}} \frac{CH}{R^{18}}$, $-CH - (CH_2)_{0-4} \frac{CH}{R^{17}} \frac{CH}{R^{18}}$, $-CH - (CH_2)_{0-4} \frac{CH}{R^{16}}$, $-CH - (CH_2)_{0-6} \frac{CH}{R^{16}}$, $-CH - (CH_2)_{0-6}$

∸C I R −C

5

$$-\frac{\text{CH}-(\text{CH}_2)_{0-4}}{\text{K}^{1}}\text{S}-(\text{CH}_2)_{0-4}} \stackrel{\text{R}^{21}}{\text{N}}-(\text{CH}_2)_{0-4}} \stackrel{\text{R}^$$

<u></u>20

8. A compound of claim 1, wherein
$$R^A$$
 is R^b when W is R^{22} R^{23} R^{25} R^{26} .

25

9. A compound of claim 4 which is (4R)-3-[8-(ethoxy-carbonyl)octanoyl]-2-(3-nitrophenyl)-4-thiazolidinecarboxylic acid.

- 1 10. A compound of claim 4 which is (4R,4'R)-3,3'-(nonane-dioyl)bis[2-(3-nitrophenyl)-4-thiazolidinecarboxylic acid methylester].
- 11. A compound according to claim 4 which is (4R)-3-(11-carboxyundecanoy1)-2-(3-cyanopheny1)-4-thiazolidinecarboxylic acid:

 (4R,4'R)-3,3'-(decaredicy1)bis[2-(3-cyanopheny1)-4-thiazolidine-carboxylic acid];

 (4R,4'R)-3,3'-(decaredicy1)bis[2-(3-cyanopheny1)-4-thiazolidine-thiazolidinecarboxylic acid;

 (4R)-3-(8-carboxyoctanoy1)-2-(3-nitropheny1)-4-thiazolidine-carboxylic acid;

 (4R,4'R)-3,3'-(nonanedicy1)bis[2-(3-nitropheny1)-4-thiazolidinecarboxylic acid];

 (4R)-3-(7-carboxyheptanoy1)-2-(2-hydroxypheny1)-4-thiazolidine-
- 12. A compound according to claim 4 which is (4R)-3
 [[(1-carboxy-3-phenylpropyl)amino]acetyl]-2-(2-hydroxyphenyl)4-thiazolidinecarboxylic acid;

 (4R)-3-[[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]acetyl]-2
 (2-hydroxyphenyl)-4-thiazolidinecarboxylic acid.

carboxylic acid.

- 13. A compound according to claim 4 which is 1-[[(1-carboxy-3-phenylpropyl)amino]acetyl]-2-(2-hydroxyphenyl)-5-pyrrolidinecarboxylic acid;
 1-[[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]acetyl]-2-(2-hydroxyphenyl)-5-pyrrolidinecarboxylic acid.
- 14. A compound of claim 4 which is (4R)-3-[[(1-carboxy3-phenylpropyl) thio]acetyl]-2-(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid.
 - 15. A compound of claim 4 which is (4R)-3-(4-carboxy-butancyl)-2-(2-hydroxyphenyl)-4-thiazolidinecarboxylic acid.

16. A process for preparing a compound of the formula [I]

$$\begin{array}{c|c}
 & Q^{1-Q^{2}} \\
 & Q^{1-Q^{2}$$

wherein

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 Q^1 and Q^2 each is metrylene or sulfur, but Q^1 and Q^2 are not sulfur at the same time;

RA is Ra or Rb;

RB and RC each is RC;

15 W is $\begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}_{L} \begin{bmatrix} R^3 \\ R^4 \end{bmatrix}_{R} \begin{bmatrix} R^5 \\ R^6 \end{bmatrix}_{R} \begin{bmatrix} R^7 \\ C \\ R^8 \end{bmatrix}_{P} \underbrace{\begin{bmatrix} R^9 \\ C \\ R^{10} \end{bmatrix}_{R} \begin{bmatrix} R^{11} \\ R^{12} \end{bmatrix}_{L} \underbrace{\begin{bmatrix} R^{13} \\ C \\ R^{14} \end{bmatrix}_{R} \begin{bmatrix} R^{15} \\ R^{16} \end{bmatrix}_{L}}_{R} \text{ wh}}_{R}$ X, Y and Z each is single bond, $-CH_2$ -, -C=C-, -C-, -C-,

or -N-|21

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1, m, n, p, q, r, s and t each is 0, 1, 2 or 3; R^1 , R^2 , R^3 , ..., R^{20} and R^{21} each is R^d ;

 R^{A} is R^{b} when W is -CH-NH-C- or -CH-(CH), wherein R^{22} , R^{23} , R^{24} , R^{25} and R^{26} each is R^{d} .

R^a is selected from the group consisting of

(i) hydrogen, lower alkyl and lower alkenyl, and

(ii) lower alkyl and lower alkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

R^b is selected from the group consisting of

(a) (i) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl, and

(ii) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl

substituted by at least one substituent selected from the group

consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl,

hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen,

nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino,

mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxy
carbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower

alkylaminosulfonyl and lower alkylsulfinyl, and

(iii) carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxy
carbonyl and heteroaryloxycarbonyl;

(b) (i) phenyl and naphthyl, and

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- (ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
- (c) (i) furyl, thienyl and pyridyl, and
 (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl,

lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy aralkyloxy, aryloxy, acyloxy, halogen, nitrocyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

R^C is selected from the group consisting of

(a) (i) hydroxy, lower alkoxy and amino, and

(ii) lower alkoxy and amino substituted by at least one substituer selected from the group consisting of lower alkyl, aralkyl, heteroarakyl, aralkenyl, heteroarakyl, hydroxy, lower alkoxy, aralkyloxy, heteroarakyloxy, aryloxy, heteroaryloxy, acyloxy, aryl, heteroaryl, substituted aralkyl and substituted aryl wherein the substituent is lower alkyl, lower alkoxy, halogen or amino;

(b) (i) aryloxy and heteroaryloxy, and(ii) aryloxy and heteroaryloxy substituted by at least one substituent selected from the group consisting of lower alkyl, hydroxy, lower alkoxy, halogen and amino, and

(c) CC-R^B;

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R^d is selected from the group consisting of

(a) (i) hydrogen, lower alkyl, lower alkenyl, aralkyl, hetero-

aralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo, and

(ii) lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, lower alkoxy, lower alkanoyl, aryl, heteroaryl, acyloxy, aroyl, hydroxy, carboxy, amino, quanidino, mercapto,

- acylamino, acylmercapto, lower alkoxycarbonyl, sulfo, halogen, nitro, cyano, sulfamoyl, lower alkylaminosulfonyl, lower alkylthio and lower alkylsulfinyl;
 - (b)(i) phenyl and naphthyl, and
- (ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
 - (c)(i) furyl, thienyl and pyridyl, and
- (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylene-dioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkyl-aminosulfonyl and lower alkylsulfinyl;

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and salts thereof

which comprises

(i) reacting a compound of the formula [II]

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$$\mathbb{R}^{A} \xrightarrow{\mathbb{Q}^{\frac{1}{2}} \mathbb{Q}^{2}}_{\mathbb{N}} = \mathbb{Q}^{2}$$
[II]

wherein R^A and R^B may include suitable protection of any reactive groups with the reactive derivative of a carboxylic acid of the formula [III] (e.g., acyl halide, acid anhydride, mixed anhydride, active ester, etc.)

HOOC-W-CO-RC

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[III]

wherein R^C and W may include suitable protection of any reactive groups, followed by removal of protective groups, if necessary to yield a compound of the formula [I];

(ii) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [IV]

$$HOOC-W^1-L$$
 [IV]

wherein $W^{\frac{1}{2}}$ is $\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix} \begin{bmatrix} 3 \\ 1 \\ 1 \end{bmatrix}$, and may include suitable protection of $\begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix} \begin{bmatrix} 3 \\ 1 \\ 1 \end{bmatrix}$.

any reactive groups, and L is a leaving group to yield a compof the formula [V]

and then reacting a compound of the formula [V] with a compound of the formula [VI]

(H)
$$X-W^2-Y-W^3-Z-W^4-CO-R^C$$
 [VI]

wherein W^2 is $\begin{bmatrix} R^5 \\ C \\ 16 \end{bmatrix}$, W^3 is $\begin{bmatrix} R^9 \\ C \\ 18 \end{bmatrix}$, W^4 is $\begin{bmatrix} R^{13} \\ C \\ 110 \end{bmatrix}$, $\begin{bmatrix} R^{15} \\ C \\ 112 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{13} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, wherein W^2 is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$, where $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$ is $\begin{bmatrix} R^{15} \\ C \\ 114 \end{bmatrix}$.

and w^2 , w^3 , w^4 , x, y, z and R^C may include suitable protection

of any reactive groups, followed by removal of protective groups, if necessary, to yield a compound of the formula [I];

(iii) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [VII]

 $HOOC-W^1-X-W^2-L$ [VII],

and then with a compound of the formula [VIII]

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$$(H) Y-V^3-Z-W^4-CO-R^C$$
 [VIII]

by the same method as (ii) above to yield a compound of the formula [I];

(iv) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [IX]

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$$100C-W^{1}-X-W^{2}-Y-W^{3}-L$$
 [IX],

and then with a compound of the formula [X]

$$(H) Z-W^{4}-CO-R^{C}$$
 [X]

by the same method as (ii) above to yield a compound of the formula [I];

(v) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [XI]

$$HOOC-W^1-X(H)$$
 [XI],

and then with a compound of the formula [XII]

[XII]

 $L-w^2-y-w^3-z-w^4-co-R^C$

by the same method as (ii) above to yield a compound of the formula [I];

5 (vi) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [XIII]

$$HOOC-W^1-X-W^2-Y(H)$$
 [XIII],

and then with a compound of the formula [XIV]

$$L-W^3-Z-W^4-CO-R^C$$
 [XIV]

by the same method as (ii) above to yield a compound of the formula [I], or

(vii) reacting a compound of the formula [II] with the reactive derivative of carboxylic acid of the formula [XV]

$$HOOC-W^{1}-X-W^{2}-Y-W^{3}-Z(H)$$
 [XV],

and then with a compound of the formula [XVI]

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$$L-W^4-CO-R^C$$
 [XVI]

by the same method as (ii) above to yield a compound of the formula [I];

furthermore converting R^B, R^C, X, Y and Z to other functional groups by the general methods, if desired, to obtain a desired compound of the formula [I].

17. A composition comprising a compound of the formula [I]

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$$R^{A} \xrightarrow{Q^{1}-Q^{2}} CO-R^{B}$$

$$CO-W-CO-R^{C}$$
[1]

wherein

 Q^1 and Q^2 each is methylene or sulfur, but Q^1 and Q^2 are not sulfur at the same time;

R^A is R^a or R^b;
R^B and R^C each is R^C;

W is $\begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}_z \begin{bmatrix} R^3 \\ C \\ R^4 \end{bmatrix}_m \times \begin{bmatrix} R^5 \\ C \\ R^6 \end{bmatrix}_n \begin{bmatrix} R^7 \\ C \\ R^9 \end{bmatrix}_z \begin{bmatrix} R^1 \\ C \\ R^1 \end{bmatrix}_z \begin{bmatrix} R^1 \\ C \\ R^1 \end{bmatrix}_z \begin{bmatrix} R^1 \\ C \\ R^1 \end{bmatrix}_z$, where:

X, Y and Z each is single bond, $-CH_2$, -C = C, -C = C, -C = C,

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$$-N$$
 N- or $-N$;

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1, m, n, p, q, r, s and t each is 0, 1, 2 or 3; R^1 , R^2 , R^3 , ..., R^{20} and R^{21} each is R^d ;

R^a is selected from the group consisting of

(i) hydrogen, lower alkyl and lower alkenyl, and

(ii) lower alkyl and lower alkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylamino-f sulfonyl and lower alkylsulfinyl;

R^b is selected from the group consisting of

(a) (i) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl, a:

(ii) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogenolower alkyl, hydroxy, lower alkoxy, halogenolower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino. dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl, and

(iii) carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl and heteroaryloxycarbonyl;

(b) (i) phenyl and naphthyl, and
(ii) phenyl and naphthyl substituted by at least one subsituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkexy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino,

acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl; (c)(i) furyl, thienyl and pyridyl, and

(ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonvl, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

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R^C is selected from the group consisting of

(a) (i) hydroxy, lower alkoxy and amino, and

(ii) lower alkoxy and amino substituted by at least one substituent selected from the group consisting of lower alkyl, aralkyl, heteroaralkyl, aralkenyl, heteroaralkyl, heteroaralkyl, hydroxy, lower alkoxy, aralkyloxy, heteroaralkyloxy, aryloxy, heteroaryloxy, acyloxy, aryl, heteroaryl, substituted aralkyl and substituted aryl wherein the substituent is lower alkyl, lower alkoxy, halogen, or amino;

(b) (i) aryloxy and heteroaryloxy, and(ii) aryloxy and heteroaryloxy substitution

(ii) aryloxy and heteroaryloxy substituted by at least one subsituent selected from the group consisting of lower alkyl, hydroxy, lower alkoxy, halogen and amino, and

(c)
$$C^{1-Q^{2}}$$
 $R^{A} \longrightarrow CO-R^{B}$

R^d is selected from the group consisting of

(a) (i) hydrogen, lower alkyl, lower alkenyl, aralkyl,
heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl,
hydroxy, carboxy, amino, mercapto and sulfo, and

- (a) (ii) lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, lower alkoxy, lower alkanoyl, aryl, heteroaryl, acyloxy, aroyl, hydroxy, carboxy, amino, guanidino,
- mercapto, acyloxy, aroyl, nydroxy, carboxy, amino, guanid mercapto, acylamino, acylmercapto, lower alkoxycarbonyl, sulfo, halogen, nitro, cyano, sulfamoyl, lower alkylaminosulfonyl, lower alkylthio and lower alkylsulfinyl;
 - (b) (i) phenyl and naphthyl, and
 - (ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl,
- lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylemercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
 - (c) (i) furyl, thienyl and pyridyl, and
- (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;

- or salts thereof in an amount sufficient to prevent or relieve diabetes mellitus associated complications consisting of cataracts, neuropathy, nephropathy and retinopathy, and pharmaceutically acceptable excipient(s).
- 18. A composition comprising a compound of the formula [I]

, wherein R^{22} ,

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$$R^{A} \xrightarrow{Q^{1} Q^{2}} CO - R^{B}$$

$$CO - W - CO - R^{C}$$
[I]

wherein Q^1 and Q^2 each is methylene or sulfur, but Q^1 and Q^2 are not sulfur at the same time;

RA is Ra or Rb;

RB and RC each is RC;

X, Y and Z each is single bond, $-CH_2^-$, $-C = C^-$

$$\begin{array}{c} & & & \\ & &$$

$$-N$$
 N- or $-N$;

25 R^1 , R^2 , R^3 , ..., R^{20} and R^{21} each is R^d ; R^A is R^b when W is

 R^{23} , R^{24} , R^{25} and R^{26} each is R^{d} ;

R^a is selected from the group consisting of

(i) hydrogen, lower alkyl and lower alkenyl, and

(ii) lower alkyl and lower alkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylamino-sulfonyl and lower alkylsulfinyl;

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R^b is selected from the group consisting of

(a) (i) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl,

(ii) aralkyl, heteroaralkyl, aralkenyl and heteroaralkenyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogenolower alkyl, hydroxy, lower alkoxy, halogenolower alkyl, hydroxy, lower alkoxy, halogenolower alkylamino, acylamino, lower alkylamino-sulfonyl, aryloxycarbonyl, sulfamoyl, lower alkylamino-sulfonyl and lower alkylsulfinyl, and

(iii) carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl and heteroaryloxycarbonyl;

(b) (i) phenyl and naphthyl, and

(ii) phenyl and naphthyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy, halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen, nitro, cyano, amino, lower alkylamino, dialkylamino, acylamino, mercapto, acylmercapto, lower alkylthio, carboxy, lower alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl, lower alkylsulfonyl and lower alkylsulfinyl;

(c) (i) furyl, thienyl and pyridyl, and
 (ii) furyl, thienyl and pyridyl substituted by at least one
substituent selected from the group consisting of lower alkyl,
lower alkenyl, halogeno-lower alkyl, hydroxy, lower alkoxy,
halogeno-lower alkoxy, aralkyloxy, aryloxy, acyloxy, halogen,
nitro, cyano, amino, lower alkylamio, dialkylamino, acylamino,
mercapto, acylmercapto, lower alkylthio, carboxy, lower
alkoxycarbonyl, aralkyloxycarbonyl, aryloxycarbonyl, sulfamoyl,
lower alkylsulfonyl, and lower alkylsulfinyl;

R^C is selected from the group consisting of

(a) (i) hydroxy, lower alkoxy and amino, and

(ii) lower alkoxy and amino substituted by at least one substituent selected from the group consisting of lower alkyl, aralkyl, heteroaralkyl, aralkenyl, heteroaralkenyl, hydroxy, lower alkoxy, aralkyloxy, heteroaralkyloxy, aryloxy, heteroaryloxy, acyloxy, aryl, heteroaryl, substituted aralkyl and substituted aryl wherein the substitutent is lower alkyl, lower alkoxy, halogen or amino;

(b) (i) aryloxy and heteroaryloxy, and
(ii) aryloxy and heteroaryloxy substituted by at least one substituent selected from the group consisting of lower alkyl, hydroxy, lower alkoxy, halogen and amino, and

 $R^{A} \xrightarrow{Q^{1}-Q^{2}} CO-R^{B}$

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Rd is selected from the group consisting of (a)(i) hydrogen, lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylallanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo, and (ii) lower alkyl, lower alkenyl, aralkyl, heteroaralkyl, alkanoyl, arylalkanoyl, heteroarylalkanoyl, hydroxy, carboxy, amino, mercapto and sulfo substituted by at least

- one substituent selected from the group consisting of lower alkyl, lower alkenyl, lower alkoxy, lower alkanoyl, aryl, heteroaryl, acyloxy, aroyl, hydroxy, carboxy, amino, quanidino, mercapto, acylamino, acylmercapto, lower alkoxycarbonyl, sulfo, halogen, nitro, cyano, sulfamoyl, lower alkylaminosu¹fonyl, lower alkylthio and lower alkylsulfinyl;
 - (b) (i) phenyl and naphthyl, and

 (ii) phenyl and naphthyl substituted by at least one
 substituent selected from the group consisting of lower
 alkyl, lower alkoxy, lower alkanoyl, acyloxy, hydroxy,
 cabroxy, amino, halogen, nitro, cyano, acylamino, mercarto,
 acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy,
 lower alkylenedioxy, lower alkoxycarbonyl, sulfo, sulfamoyl,
 lower alkylaminosulfonyl and lower alkylsulfinyl;
 - (c) (i) furyl, thienyl and pyridyl, and
 (ii) furyl, thienyl and pyridyl substituted by at least one substituent selected from the group consisting of lower alkyl, lower alkoxy, lower alkanoyl, acyloxy,
- hydroxy, carboxy, amino, halogen, nitro, cyano, acylamino, mercapto, acylmercapto, halogeno-lower alkyl, halogeno-lower alkoxy, halogeno-lower alkoxycarbonyl, sulfo, sulfamoyl, lower alkylaminosulfonyl and lower alkylsulfinyl;
- or salts thereof in an amount sufficient to reduce blood pressure and pharmaceutically acceptable excipient(s).
 - 19. A compound according to claim 1 to 16 for use in a method for therapy or prophylaxis.
 - 20. Use of a compound according to claim 1 to 16 in a process for producing pharmaceutical compositions.

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EUROPEAN SEARCH REPORT

Application number

EP 80 10 7869

	OCUMENTS CONSIDERED TO BE RELEVANT		•	CLASSIFICATION OF THE APPLICATION (Int. Cl.)	
Category	Citation of document with indic passages	ation, where appropriate, of relevant	Relevant to claim	(inc Oil)	
*	US - A - 4 154 et al.) * Columns 1-2 *	937 (D.W. CUSHMAN	1-3,5 6,7, 16	C 07 D 277/0 207/1 A 61 K 31/4	
*	GB - A - 2 000 PHARM. LTD.) * Pages 1-2 *	508 (YOSHITGMI	1 - 5, 7,16	31/4	
		204 (SANDOZ S.A.)	1 - 5 _.	TECHNICAL FIELDS SEARCHED (Int. CI. ²)	
	FR - A - 2 412 ET CIE) *"Revendication	537 (SCIENCE UNION	1,2	C 07 D 277/0 277/	
	FR - A - 2 340 AND SONS) *"Revendication	933 (E.R.SQUIBB	1-3, 5-7	·	
	FR - A - 2 340 AND SONS) *"Revendication	932 (E.R SQUIBB		CATEGORY OF CITED DOCUMENTS X: particularly relevant A: technological background	
	FR - A - 2 023 *"Revendication	741 (EPROVA AG)	1	O: non-written disclosure P: intermediate document T: theory or principle underlying the invention E: conflicting application	
P	EP - A - 0 007 PHARM) *"Revendication	*	1-5	D: document cited in the application L: citation for other reasons	
0		t has been drawn up for all claims		 member of the same patent family, corresponding document 	
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EUROPEAN SEARCH REPORT

Application number

EP 80 10 7869 -2-

	DOCUMENTS CONSIDERED TO BE RELEVANT		CLASSIFICATION OF THE APPLICATION (Int. Cl. 3)
Category	Citation of document with Indication, where appropriate, of relevant passages	Relevant to claim	The second of th
P	FR - A - 2 445 324 (SANTEN PHARM) *"Revendications"*	1-5	•
P	FR - A - 2 440 365 (SANTEN PHARM)	1-5	
	"Revendications"		
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F	FR - A - 2 434 150 (YOSHITOMI PHARM.)	1-5	TECHNICAL CITY OF
	"Revendications"		TECHNICAL FIELDS SEARCHED (Int. Ci. ²)
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